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=> file biosis medline caplus wpids uspatfull
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                                                   ENTRY
                                                            SESSION
FULL ESTIMATED COST
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                                                               0.22
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CA INDEXING COPYRIGHT (C) 2010 AMERICAN CHEMICAL SOCIETY (ACS)
*** YOU HAVE NEW MAIL ***
=> s (nucleotide or nucleoside) and base (4a) (label? or marker or dve)
L1
         4804 (NUCLEOTIDE OR NUCLEOSIDE) AND BASE (4A) (LABEL? OR MARKER OR
              DYE)
=> s l1 and linker (4a) (label? or marker or dye)
          531 L1 AND LINKER (4A) (LABEL? OR MARKER OR DYE)
=> s 12 and linker (6a) polymer
L3
           27 L2 AND LINKER (6A) POLYMER
=> s 13 and water soluble polymer
            0 L3 AND WATER SOLUBLE POLYMER
L4
=> s 13 and linker (4a) 50
L5
            8 L3 AND LINKER (4A) 50
=> dup rem 15
PROCESSING COMPLETED FOR L5
             8 DUP REM L5 (0 DUPLICATES REMOVED)
1.6
=> d 16 bib abs 1-8
     ANSWER 1 OF 8 USPATFULL on STN
L6
       2007:237090 USPATFULL
AN
       Compositions for the electronic detection of analytes utilizing
      monolavers
      Kayyem, Jon Faiz, Pasadena, CA, UNITED STATES
       O'Connor, Stephen D., Pasadena, CA, UNITED STATES
      Clinical Micro Sensors, Inc. (U.S. corporation)
PΤ
      US 20070207465
                         A1 20070906
      US 7393645
                         B2 20080701
AΤ
      US 2005-208384
                        A1 20050819 (11)
RLT
      Division of Ser. No. US 1999-452277, filed on 30 Nov 1999, GRANTED, Pat.
      No. US 7160678 Continuation of Ser. No. US 1997-911085, filed on 14 Aug
```

```
1997, GRANTED, Pat. No. US 6090933 Continuation of Ser. No. US
       1997-873978, filed on 12 Jun 1997, GRANTED, Pat. No. US 7014992
       Continuation of Ser. No. US 1996-743798, filed on 5 Nov 1996, GRANTED,
       Pat. No. US 6096273 Continuation of Ser. No. US 1997-911589, filed on 14
       Aug 1997, GRANTED, Pat. No. US 6232062 Continuation of Ser. No. US
       1997-873597, filed on 12 Jun 1997, PENDING
PRAT
       WO 1997-US20014
                               19971105
       US 1997-40155P
                               19970307 (60)
       US 1997-49489P
                               19970612 (60)
       US 1997-40153P
                              19970307 (60)
DT
      Utility
FS
      APPLICATION
LREP
      MORGAN, LEWIS & BOCKIUS, LLP, ONE MARKET SPEAR STREET TOWER, SAN
       FRANCISCO, CA, 94105, US
CLMN
      Number of Claims: 24
ECI.
      Exemplary Claim: 1-21
DRWN
      41 Drawing Page(s)
LN.CNT 4693
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AB
       The present invention relates to the use of self-assembled monolayers
       with mixtures of conductive oligomers and insulators to detect target
       analytes.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
     ANSWER 2 OF 8 USPATFULL on STN
1.6
       2007:7693 USPATFULL
AN
TI
       Compositions for the electronic detection of analytes utilizing
       monolavers
ΤN
       Kayyem, Jon Faiz, South Pasadena, CA, UNITED STATES
       O'Connor, Stephen D., Pasadena, CA, UNITED STATES
       Clinical Micro Sensors, Inc., Pasadena, CA, UNITED STATES (U.S.
PA
       corporation)
PΙ
       US 7160678
                           B1 20070109
      US 1999-452277
AΙ
                               19991130 (9)
RLI
       Continuation of Ser. No. US 1997-911085, filed on 14 Aug 1997, Pat. No.
       US 6090933 Continuation of Ser. No. US 1997-911589, filed on 14 Aug
       1997, Pat. No. US 6232062 Continuation of Ser. No. US 1997-873978, filed
      on 12 Jun 1997, PENDING Continuation of Ser. No. US 1997-873597, filed
      on 12 Jun 1997, PENDING Continuation of Ser. No. US 1996-743798, filed
      on 5 Nov 1996, Pat. No. US 6096273
PRAI
      WO 1997-US20014
                              19971105
      US 1998-73014P
                              19980129 (60)
       US 1997-49489P
                              19970612 (60)
       US 1997-40153P
                              19970307 (60)
       US 1997-40155P
                              19970307 (60)
      Utility
FS
      GRANTED
EXNAM Primary Examiner: Whisenant, Ethan
LREP
       Dorsey & Whitney LLP, Silva, Robin M.
CLMN
      Number of Claims: 20
ECL
       Exemplary Claim: 1
DRWN
      93 Drawing Figure(s); 41 Drawing Page(s)
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
```

The present invention relates to the use of self-assembled monolayers with mixtures of conductive oligomers and insulators to detect target

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

analytes.

```
1.6
     ANSWER 3 OF 8 USPATFULL on STN
AN
       2003:237907 USPATFULL
TT
       Compositions and methods for the therapy and diagnosis of colon cancer
TN
       King, Gordon E., Shoreline, WA, UNITED STATES
       Meagher, Madeleine Joy, Seattle, WA, UNITED STATES
       Xu, Jiangchun, Bellevue, WA, UNITED STATES
       Secrist, Heather, Seattle, WA, UNITED STATES
       Jiang, Yugiu, Kent, WA, UNITED STATES
PA
       Corixa Corporation, Seattle, WA, UNITED STATES, 98104 (U.S. corporation)
PΙ
       US 20030166064
                           A1 20030904
ΑI
       US 2002-99926
                           A1 20020314 (10)
       Continuation-in-part of Ser. No. US 2001-33528, filed on 26 Dec 2001,
RLI
       PENDING Continuation-in-part of Ser. No. US 2001-920300, filed on 31 Jul
       2001, PENDING
PRAI
       US 2001-302051P
                               20010629 (60)
       US 2001-279763P
                               20010328 (60)
       US 2000-223283P
                              20000803 (60)
DT
       Utility
FS
       APPLICATION
LREP
       SEED INTELLECTUAL PROPERTY LAW GROUP PLLC, 701 FIFTH AVE, SUITE 6300,
       SEATTLE, WA, 98104-7092
CLMN
       Number of Claims: 17
ECL
       Exemplary Claim: 1
DRWN
      No Drawings
LN.CNT 8531
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       Compositions and methods for the therapy and diagnosis of cancer,
       particularly colon cancer, are disclosed. Illustrative compositions
       comprise one or more colon tumor polypeptides, immunogenic portions
       thereof, polynucleotides that encode such polypeptides, antigen
       presenting cell that expresses such polypeptides, and T cells that are
       specific for cells expressing such polypeptides. The disclosed
       compositions are useful, for example, in the diagnosis, prevention
       and/or treatment of diseases, particularly colon cancer.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L6
     ANSWER 4 OF 8 USPATFULL on STN
ΑN
       2003:106233 USPATFULL
ΤТ
       Compositions and methods for the therapy and diagnosis of pancreatic
IN
       Benson, Darin R., Seattle, WA, UNITED STATES
       Kalos, Michael D., Seattle, WA, UNITED STATES
       Lodes, Michael J., Seattle, WA, UNITED STATES
       Persing, David H., Redmond, WA, UNITED STATES
       Hepler, William T., Seattle, WA, UNITED STATES
       Jiang, Yugiu, Kent, WA, UNITED STATES
       Corixa Corporation, Seattle, WA, UNITED STATES, 98104 (U.S. corporation)
PA
ΡI
                          A1 20030417
       US 20030073144
AΙ
       US 2002-60036
                           A1 20020130 (10)
PRAI
       US 2001-333626P
                               20011127 (60)
                               20010712 (60)
       US 2001-305484P
       US 2001-265305P
                               20010130 (60)
       US 2001-267568P
                               20010209 (60)
       US 2001-313999P
                               20010820 (60)
       US 2001-291631P
                               20010516 (60)
       US 2001-287112P
                               20010428 (60)
       US 2001-278651P
                               20010321 (60)
       US 2001-265682P
                              20010131 (60)
      Utility
FS
      APPLICATION
```

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LREP
       SEED INTELLECTUAL PROPERTY LAW GROUP PLLC, 701 FIFTH AVE, SUITE 6300,
       SEATTLE, WA, 98104-7092
      Number of Claims: 17
CLMN
      Exemplary Claim: 1
ECI.
DRWN No Drawings
LN.CNT 14253
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       Compositions and methods for the therapy and diagnosis of cancer,
       particularly pancreatic cancer, are disclosed. Illustrative compositions
       comprise one or more pancreatic tumor polypeptides, immunogenic portions
       thereof, polynucleotides that encode such polypeptides, antigen
       presenting cell that expresses such polypeptides, and T cells that are
       specific for cells expressing such polypeptides. The disclosed
       compositions are useful, for example, in the diagnosis, prevention
       and/or treatment of diseases, particularly pancreatic cancer.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L6
    ANSWER 5 OF 8 USPATFULL on STN
ΑN
       2003:203373 USPATFULL
ΤI
       Electronic methods for the detection of analytes utilizing monolayers
IN
       Yu, Changjun, Pasadena, CA, United States
      Clinical Micro Sensors, Inc., Pasadena, CA, United States (U.S.
PA
      corporation)
       US 6600026
                          B1 20030729
      US 1999-306653
                              19990506 (9)
AΤ
RLT
      Continuation of Ser. No. US 1998-135183, filed on 17 Aug 1998
      US 1998-84652P
                              19980506 (60)
PRAI
      US 1998-84509P
                              19980506 (60)
DТ
      Utility
FS
      GRANTED
EXNAM Primary Examiner: Riley, Jezia
LREP
      Silva, Robin M., Kosslak, Renee M., Dorsey & Whitney, LLP
CLMN
     Number of Claims: 12
ECL
      Exemplary Claim: 1
DRWN
     93 Drawing Figure(s); 41 Drawing Page(s)
LN.CNT 4573
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       The present invention relates to the use of self-assembled monolayers
       with mixtures of conductive oligomers and insulators to detect target
       analytes.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
1.6
     ANSWER 6 OF 8 USPATFULL on STN
AN
       2002:272801 USPATFULL
ΤТ
       Compositions and methods for the therapy and diagnosis of colon cancer
       Stolk, John A., Bothell, WA, UNITED STATES
TN
       Xu, Jiangchun, Bellevue, WA, UNITED STATES
       Chenault, Ruth A., Seattle, WA, UNITED STATES
       Meagher, Madeleine Joy, Seattle, WA, UNITED STATES
       Corixa Corporation, Seattle, WA, UNITED STATES, 98104 (U.S. corporation)
PA
ΡI
      US 20020150922
                          A1 20021017
      US 2001-998598
ΑI
                          A1 20011116 (9)
      US 2001-304037P
PRAI
                               20010710 (60)
       US 2001-279670P
                              20010328 (60)
      US 2001-267011P
                              20010206 (60)
      US 2000-252222P
                              20001120 (60)
DT
      Utility
FS
      APPLICATION
      SEED INTELLECTUAL PROPERTY LAW GROUP PLLC, 701 FIFTH AVE, SUITE 6300,
```

SEATTLE, WA, 98104-7092 Number of Claims: 17 CLMN ECI. Exemplary Claim: 1 DRWN No Drawings LN.CNT 9233 CAS INDEXING IS AVAILABLE FOR THIS PATENT. Compositions and methods for the therapy and diagnosis of cancer, particularly colon cancer, are disclosed. Illustrative compositions comprise one or more colon tumor polypeptides, immunogenic portions thereof, polynucleotides that encode such polypeptides, antigen presenting cell that expresses such polypeptides, and T cells that are specific for cells expressing such polypeptides. The disclosed compositions are useful, for example, in the diagnosis, prevention and/or treatment of diseases, particularly colon cancer. CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 7 OF 8 USPATFULL on STN AN 2002:243051 USPATFULL ΤI Compositions and methods for the therapy and diagnosis of ovarian cancer TN Algate, Paul A., Issaquah, WA, UNITED STATES Jones, Robert, Seattle, WA, UNITED STATES Harlocker, Susan L., Seattle, WA, UNITED STATES Corixa Corporation, Seattle, WA, UNITED STATES, 98104 (U.S. corporation) PA A1 20020919 PΙ US 20020132237 US 2001-867701 A1 20010529 (9) AΤ PRAT US 2000-207484P 20000526 (60) Utility DT FS APPLICATION

CLMN Number of Claims: 11 ECL Exemplary Claim: 1

LREP

IN

DRWN No Drawings

LN.CNT 25718

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

SEATTLE, WA, 98104-7092

AB Compositions and methods for the therapy and diagnosis of cancer, particularly ovarian cancer, are disclosed. Illustrative compositions comprise one or more ovarian tumor polypeptides, immunogenic portions thereof, polynucleotides that encode such polypeptides, antigen presenting cell that expresses such polypeptides, and T cells that are specific for cells expressing such polypeptides. The disclosed compositions are useful, for example, in the diagnosis, prevention and/or treatment of diseases, particularly ovarian cancer.

SEED INTELLECTUAL PROPERTY LAW GROUP PLLC. 701 FIFTH AVE. SUITE 6300.

## CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 8 OF 8 USPATFULL on STN L6

2002:242791 USPATFULL AN

TΙ Compositions and methods for the therapy and diagnosis of colon cancer

King, Gordon E., Shoreline, WA, UNITED STATES Meagher, Madeleine Joy, Seattle, WA, UNITED STATES Xu, Jiangchun, Bellevue, WA, UNITED STATES

Secrist, Heather, Seattle, WA, UNITED STATES Corixa Corporation, Seattle, WA, UNITED STATES (U.S. corporation)

PA PΙ US 20020131971

A1 20020919 AΤ US 2001-33528 A1 20011226 (10)

RLT Continuation-in-part of Ser. No. US 2001-920300, filed on 31 Jul 2001,

PENDING

PRAI US 2001-302051P 20010629 (60) US 2001-279763P 20010328 (60) US 2000-223283P 20000803 (60)

DT Utility

FS APPLICATION

LREP SEED INTELLECTUAL PROPERTY LAW GROUP PLLC, 701 FIFTH AVE, SUITE 6300, SEATTLE, WA. 98104-7092

CLMN Number of Claims: 17

ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 8083

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Compositions and methods for the therapy and diagnosis of cancer, particularly colon cancer, are disclosed. Illustrative compositions comprise one or more colon tumor polypeptides, immunogenic portions thereof, polynucleotides that encode such polypeptides, antigen presenting cell that expresses such polypeptides, and T cells that are specific for cells expressing such polypeptides. The disclosed compositions are useful, for example, in the diagnosis, prevention and/or treatment of diseases, particularly colon cancer.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

=> d 16 1 kwic

L6 ANSWER 1 OF 8 USPATFULL on STN

DRWD

. . portion of the target sequence 5, a second portion 42 that hybridizes to the capture probe 10 and a recruitment linker 50 comprising ETMs 6. A similar embodiment is shown in FTG. 6E, where the label probe 40 has an additional recruitment linker 50. FTG. 6F depicts a label probe 40 comprising a first portion 41 that hybridizes to a portion of the target sequence 5 and a recruitment linker 50 with attached ETMs 6. The parentheses highlight that for any particular target sequence 5 more

than one label probe 40. . . . DRWD . . . portion 41 of the label probe 40 can hybridize to all (FIG.

6R) or part (FIG. 6Q) of the recruitment linker 50.

DRWD FIG. 11 depicts the synthetic scheme of a preferred attachment of an

ETM. in this case ferrocene, to a nucleoside via the

DRWD FIG. 13 depicts the synthesis of an insulator to the ribose of a nucleoside for attachment to an electrode.

DRWD . . . depicts a schematic of the synthesis of simultaneous incorporation of multiple ETMs into a nucleic acid, using a "branch" point nucleoside.

DRWD ... and attachments of ETMs. In FIGS. 21A-C, the recruitment linker is nucleic acid; in FIGS. 21D and E, is not. A=nucleoside replacement; B=attachment to a base; C=attachment to a ribose; D=attachment to a phosphate; E=metallocene polymer (although as described herein, this.

DETD . al., Angew. Chem. Intl. Ed. English 30:423 (1991); Letsinger et al., J. Am. Chem. Soc. 110:4470 (1988); Letsinger et al., Nucleoside & Nucleotide 13:1597 (1994); Chapters 2 and 3, ASC Symposium Series 580, "Carbohydrate Modifications in Antisense Research", Ed. Y. S. Sanghui and.

DETD sequences, as this reduces non-specific hybridization, as is generally described in U.S. Pat. No. 5,681,702. As used herein, the term "nucleoside" includes nucleotides as well as nucleoside and nucleotide analogs, and modified nucleosides such as amino modified nucleosides. In addition, "nucleoside" includes non-naturally occurring analog structures.

Thus for example the individual units of a peptide nucleic acid, each

- containing a base, are referred to herein as a nucleoside.
- DETD . . . a conductive oligomer, as is more fully described below, the length of the conductive oligomer is such that the closest nucleotide of the nucleic acid is positioned from about 6 Å to about 100 Å (although distances of up to 500. . .
- DETD . . embodiment, the capture binding ligand is a nucleic acid, and the attachment is via attachment to the base of the nucleoside , via attachment to the backbone of the nucleic acid (either the ribose, the obsophate, or to an analogous group of.
- DETD In a preferred embodiment, the conductive oligomer is attached to the base of a nucleoside of the nucleic acid. This may be done in several ways, depending on the otigomer, as is described below. In one embodiment, the oligomer is attached to a terminal nucleoside, i.e. either the 3' or 5' nucleoside of the nucleic acid.

  Alternatively, the conductive oligomer is attached to an internal nucleoside.
- DETD . . . depicted herein may have hydrogen, hydroxy, phosphates or other groups such as amino groups attached. In addition, the pentose and nucleoside structures depicted herein are depicted non-conventionally, as mirror images of the normal rendering. In addition, the pentose and nucleoside structures may also contain additional groups, such as protecting groups, at any position, for example as needed during synthesis.
- DETD . . . it should be understood that the site of attachment in this embodiment may be to a 3' or 5' terminal nucleotide, or to an internal nucleotide, as is more fully described below.
- DETD . . row transition metals are potential candidates as redox moieties that are covalently attached to either the ribose ring or the nucleoside base of nucleic acid. Other potentially suitable organometallic ligands include cyclic arenes such as benzene, to yield bis (arene)metal compounds. .
- DETD . . . be. The solution binding ligand either directly comprises a recruitment linker that comprises at least one ETM, or the recruitment linker is part of a label probe that will bind to the solution binding ligand.
- DETD . via (1) a base; (2) the backbone, including the ribose, the phosphate, or comparable structures in nucleic acid analogs; (3) nucleoside replacement, described below, or (4) metallocene polymers, as described below. In a preferred embodiment, the recruitment linker is non-nucleic acid. . . .
- DETD . . . in a variety of positions. Preferred embodiments include, but are not limited to, (1) attachment to the base of the nucleoside , (2) attachment of the ETM as a base replacement, (3) attachment to the backbone of the nucleic acid, including either. . .
- DETD In a preferred embodiment, the ETM is attached to the base of a nucleoside as is generally outlined above for attachment of the conductive oligomer. Attachment can be to an internal nucleoside or a terminal nucleoside.
- DETD . . . defined above. Again, it will be appreciated by those in the art, a linker ("2") may be included between the nucleoside and the ETM.
- DETD In a preferred embodiment, the ETM attached to a nucleoside is a metallocene; i.e. the L and L, of Structure 31 are both metallocene ligands, Lm, as described above. Structure. . . .
- DETD . . . any position of the ribose-phosphate backbone of the nucleic acid, i.e. either the 5' or 3' terminus or any internal nucleoside. Ribose in this case can include ribose analogs. As is known in the art, nucleosides that are modified at either. . .
- DETD When the ETM is attached to the base or the backbone of the nucleoside, it is possible to attach the ETMs via "dendrimer" structures, as is more fully outlined below. As is generally depicted.

- . . terminal hydroxy groups can then be used in phosphoramidite reactions to add ETMs, as is generally done below for the nucleoside replacement and metallocene polymer reactions.
- DEID In a preferred embodiment, an ETM such as a metallocene is used as a "
  nucleoside replacement", serving as an ETM. For example, the
  distance between the two cyclopentadiene rings of ferrocene is similar
  to the.
- DETD . . . depicts metallocenes, and particularly ferrocene, this same general idea can be used to add ETMs in addition to metallocenes, as nucleoside replacements or in polymer embodiments, described below. Thus, for example, when the ETM is a transition metal complex other than.
- DEID . . nucleic acids each made up of a traditional nucleic acid or analog (nucleic acids in this case including a single nucleoside ), may be covalently attached to each other via a metallocene. Viewed differently, a metallocene derivative or substituted metallocene is provided, .
- DETD . . additional substituent groups to one or both of the aromatic rings of the metallocene (or ETM). For example, as these nucleoside replacements are generally part of probe sequences to be hybridized with a substantially complementary nucleic acid, for example a target.
- DETD . as outlined in U.S. Pat. No. 5,124,246, using modified functionalized nucleotides. The general idea is as follows. A modified phosphoramidite nucleotide is generated that can ultimately contain a free hydroxy group that can be used in the attachment of phosphoramidite ETMs. will be appreciated by those in the art, nucleic acid analogs containing other structures can also be used). The modified nucleotide is incorporated into a nucleic acid, and any hydroxy protecting groups are removed, thus leaving the free hydroxyl. Upon the.
- DETD . . this general idea is outlined in the Figures. In this embodiment, the 2' position of a ribose of a phosphoramidite nucleotide is first functionalized to contain a protected hydroxy group, in this case via an oxo-linkage, although any number of linkers can be used, as is generally described herein for Z linkers. The protected modified nucleotide is then incorporated via standard phosphoramidite chemistry into a growing nucleic acid. The
- protecting group is removed, and the free. . .

  DETD In a preferred embodiment, the recruitment linker is not nucleic acid, and instead may be any sort of linker or polymer. As will be appreciated by those in the art, generally any linker or polymer that can be modified to contain ETMs can be used.
- or polymer that can be modified to Contain Fins can be used.

  In general, the polymers or linkers should be reasonably soluble. .

  DETD In a preferred embodiment, the recruitment linker comprises a metallocene polymer, as is described above.
- DETD the solution binding ligand or the first portion of the label probe will depend on the composition of the recruitment linker and of the label and/or binding ligand, as will be appreciated
- by those in the art. When either the label probe or the binding. .

  DETD When non-nucleic acid recruitment linkers are used, attachment of the linker/polymer of the recruitment linker will be done eenerally using standard chemical techniques, such as will
- be appreciated by those in the art For example,.

  . when the target sequence itself is modified to contain a binding partner, the binding partner is attached via a modified nucleotide that can be enzymatically attached to the target sequence, for example during a PCR target amplification step.
- Alternatively, the binding.

  DETD ... oligonucleotide segments emanating from a point of origin to form a branched structure. The point of origin may be another

nucleotide segment or a multifunctional molecule to which at least three segments can be covalently or tightly bound. "Comb-like" branched amplifier. . . multiplicity of sidechain oligonucleotides extending from the backbone. In either conformation, the pendant segments will normally depend from a modified nucleotide or other organic molety having the appropriate functional groups for attachment of oligonucleotides. Furthermore, in either conformation, a large number. .

- DETD The compositions may be made in several ways. A preferred method first synthesizes a conductive oligomer attached to a nucleoside, with addition of additional nucleosides to form the capture probe followed by attachment to the electrode. Alternatively, the whole capture. . . .
- DETD . . . a preferred embodiment, the compositions of the invention are made by first forming the conductive oligomer covalently attached to the nucleoside, followed by the addition of additional nucleosides to form a capture probe nucleic acid, with the last step comprising the.
- DETD The attachment of the conductive oligomer to the nucleoside may be done in several ways. In a preferred embodiment, all or part of the conductive oligomer is synthesized first (generally with a functional group on the end for attachment to the electrode), which is then attached to the nucleoside. Additional nucleosides are then added as required, with the last step generally being attachment to the electrode. Alternatively, oligomer units are added one at a time to the nucleoside, with addition of additional nucleosides and attachment to the electrode. A number of representative syntheses are shown in the Figures.
- DETD The conductive oligomer is then attached to a nucleoside that may contain one (or more) of the oligomer units, attached as depicted herein.
- DETD Alternatively, attachment to the base may be done by making the nucleoside with one unit of the oligomer, followed by the addition of others.
- DETD . . . DNA polymerase, T7 DNA polymerase, Taq DNA polymerase, reverse transcriptase, and RNA polymerases. For the incorporation of a 3' modified nucleoside to a nucleic acid, terminal deoxynucleotidyltransferase may be used. (Ratliff, Terminal
- deoxynucleotidyltransferase. In The Enzymes, Vol 14A. P. D. Boyer. .

  DETD In a preferred embodiment, the modified nucleoside is converted to the phosphoramidite or Hphosphonate form, which are then used in solid-phase or solution syntheses of oligonucleotides. In this
- way the modified nucleoside, either for attachment at the ribose (i.e. amino- or thiol-modified nucleosides) or the base, is incorporated into the oligonucleotide at.

  DETD For attachment of a group to the 3' terminus, a preferred method
- utilizes the attachment of the modified nucleoside (or the nucleoside replacement) to controlled pore glass (CPG) or other oligomeric supports. In this embodiment, the modified nucleoside is protected at the Vend with DIVIT, and then reacted with succinic anhydride with activation. The resulting succinyl compound is.
- DETD . used as the ETM, synthesis may occur in several ways. In a preferred embodiment, the ligand(s) are added to a nucleoside, followed by the transition metal ion, and then the nucleoside with the transition metal complex attached is added to an oligonucleotide, i.e. by addition to the nucleic acid synthesizer. Alternatively.
- DETD In a preferred embodiment, ETMs are attached to a base of the nucleoside. This may be done in a variety of ways. In one embodiment, amino groups of the base, either naturally occurring.
- DETD . . . subunits, and thus additional subunits are attached to form

```
the conductive oligomer. The conductive oligomer is then attached to a
      nucleoside, and additional nucleosides attached. The protecting
      group is then removed and the sulfur-gold covalent attachment is made.
      Alternatively, all or. . . atom is added, or a sulfur atom is added
      and then protected. The conductive oligomer is then attached to a
      nucleoside, and additional nucleosides attached. Alternatively,
      the conductive oligomer attached to a nucleic acid is made, and then
      either a subunit.
DETD
       . . trimethylsilylethyl group to a sulfhydryl; 2) adding
      additional subunits to form the conductive oligomer; 3) adding at least
      a first nucleoside to the conductive oligomer; 4) adding
      additional nucleosides to the first nucleoside to form a
      nucleic acid; 5) attaching the conductive oligomer to the gold
      electrode. This may also be done in.
       . . . the PNA. By "monomeric subunit of PNA" herein is meant the
DETD
      --NH--CH.sub.2--CH.sub.2--N(COCH.sub.2-Base)-CH.sub.2--CO-monomer, or
      derivatives (herein included within the definition of "
      nucleoside") of PNA. For example, the number of carbon atoms in
      the PNA backbone may be altered; see generally Nielsen et. . .
DETD
      Synthesis of Nucleoside Modified with Ferrocene at the 2'
      Position
DETD
      Synthesis of "Branched" Nucleoside
DETD
      Synthesis of Nucleoside with Ferrocene Attached Via a
DETD
      Synthesis of Nucleoside Attached to an Insulator
DETD
       . . . sequence MT1 (SEQ ID NO:18) was added, that comprises a
      sequence complementary to D112 (SEQ ID NO:7) and a 20 base
      sequence complementary to the label probe D358 (SEQ ID NO:19)
      were combined; in this case, the label probe D358 (SEQ ID NO:19) was
      added to. . .
DETD
       . . . target sequence LP280 (SEQ ID NO:22) was added, that comprises
      a sequence complementary to the capture probe and a 20 base
      sequence complementary to the label probe D335 (SEQ ID NO:21)
      were combined; in this case, the label probe D335 (SEQ ID NO:21) was
      added to. . .
=> d his
     (FILE 'HOME' ENTERED AT 11:49:49 ON 04 JUL 2010)
    FILE 'BIOSIS, MEDLINE, CAPLUS, WPIDS, USPATFULL' ENTERED AT 11:50:31 ON
    04 JUL 2010
          4804 S (NUCLEOTIDE OR NUCLEOSIDE) AND BASE (4A) (LABEL? OR MARKER OR
           531 S L1 AND LINKER (4A) (LABEL? OR MARKER OR DYE)
            27 S L2 AND LINKER (6A) POLYMER
             0 S L3 AND WATER SOLUBLE POLYMER
             8 S L3 AND LINKER (4A) 50
             8 DUP REM L5 (0 DUPLICATES REMOVED)
=> s 13 not 16
           19 L3 NOT L6
=> dup rem 17
PROCESSING COMPLETED FOR L7
            19 DUP REM L7 (0 DUPLICATES REMOVED)
=> d 18 bib abs 1-19
L8 ANSWER 1 OF 19 USPATFULL on STN
AN
     2009:232895 USPATFULL
```

L1

1.2

1.3

1.4 L5

L6

L7

```
Alternate labeling strategies for single molecule sequencing
TN
       Korlach, Jonas, Newark, CA, UNITED STATES
       Roitman, Daniel, Menlo Park, CA, UNITED STATES
       Eid, John, San Francisco, CA, UNITED STATES
       Otto, Geoff, San Carlos, CA, UNITED STATES
       Hardenbol, Paul, San Francisco, CA, UNITED STATES
       Flusberg, Benjamin, Palo Alto, CA, UNITED STATES
       Pacific Biosciences of California, Inc., Menlo Park, CA, UNITED STATES
       (U.S. corporation)
PΙ
      US 20090208957
                          A1 20090820
AΙ
      US 2008-315626
                          A1 20081203 (12)
PRAI
      US 2007-5407P
                               20071204 (61)
DT
      Utility
FS
      APPLICATION
LREP
       QUINE INTELLECTUAL PROPERTY LAW GROUP, P.C., P O BOX 458, ALAMEDA, CA,
       94501, US
CLMN
      Number of Claims: 58
ECL
      Exemplary Claim: 1
DRWN
       1 Drawing Page(s)
LN.CNT 2173
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AB
       Systems and methods of enhancing fluorescent labeling strategies as well
       as systems and methods of using non-fluorescent and/or non-optic
       labeling strategies, e.g., as with single molecule sequencing using
       ZMWs, are described.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L8
     ANSWER 2 OF 19 USPATFULL on STN
       2006:167005 USPATFULL
AN
TI
       Chemical amplification for the synthesis of patterned arrays
IN
       Beecher, Jody E., Mountain View, CA, UNITED STATES
       Goldberg, Martin J., San Jose, CA, UNITED STATES
       McGall, Glenn H., Mountain View, CA, UNITED STATES
       Affymetrix, Inc, Santa Clara, CA, UNITED STATES (U.S. corporation)
PA
PΙ
      US 20060141511
                          A1 20060629
ΑI
      US 2005-291248
                          A1 20051201 (11)
RLI
       Continuation of Ser. No. US 2004-840841, filed on 7 May 2004, PENDING
       Continuation of Ser. No. US 2000-578282, filed on 25 May 2000, GRANTED,
       Pat. No. US 6770436 Continuation of Ser. No. US 1997-969227, filed on 13
      Nov 1997, GRANTED, Pat. No. US 6083697
PRAI
      US 1996-30826P
                               19961114 (60)
DT
      Utility
FS
      APPLICATION
LREP
      BANNER & WITCOFF LTD.,, COUNSEL FOR AFFYMETRIX, 1001 G STREET , N.W.,
       ELEVENTH FLOOR, WASHINGTON, DC, 20001-4597, US
CLMN
      Number of Claims: 17
ECL
      Exemplary Claim: 1-51
DRWN
      7 Drawing Page(s)
LN.CNT 1217
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AB
       Radiation-activated catalysts (RACs), autocatalytic reactions, and
       protective groups are employed to achieve a highly sensitive, high
       resolution, radiation directed combinatorial synthesis of pattern arrays
       of diverse polymers. When irradiated, RACs produce catalysts that can
       react with enhancers, such as those involved in autocatalytic reactions.
       The autocatalytic reactions produce at least one product that removes
       protecting groups from synthesis intermediates. This invention has a
       wide variety of applications and is particularly useful for the solid
      phase combinatorial synthesis of polymers.
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CAS INDEXING IS AVAILABLE FOR THIS PATENT.
    ANSWER 3 OF 19 USPATFULL on STN
1.8
       2006:101389 USPATFULL
AN
TT
       Whole cell engineering by mutagenizing a substantial portion of a
       starting genome, combining mutations, and optionally repeating
       Short, Jay M., Rancho Santa Fe, CA, UNITED STATES
PA
       Diversa Corporation, San Diego, CA, UNITED STATES (U.S. corporation)
ΡI
       US 7033781
                           B1 20060425
ΑI
      US 2000-677584
                               20000930 (9)
RLI
      Continuation-in-part of Ser. No. US 2000-594459, filed on 14 Jun 2000,
       PENDING Continuation-in-part of Ser. No. US 2000-552289, filed on 9 Mar
       2000, Pat. No. US 6358709 Continuation-in-part of Ser. No. US
       2000-498557, filed on 4 Feb 2000, PENDING Continuation-in-part of Ser.
       No. US 2000-495052, filed on 31 Jan 2000, PENDING
      US 1999-156815P
                              19990929 (60)
PRAT
DT
      Utility
FS
      GRANTED
EXNAM Primary Examiner: Park, Hankyel T.
LREP
       Love, Jane M., Hale and Dorr LLP
CLMN
      Number of Claims: 24
ECL
      Exemplary Claim: 1
       30 Drawing Figure(s); 28 Drawing Page(s)
DRWN
LN.CNT 36686
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
```

An invention comprising cellular transformation, directed evolution, and screening methods for creating novel transgenic organisms having desirable properties. Thus in one aspect, this invention relates to a method of generating a transgenic organism, such as a microbe or a plant, having a plurality of traits that are differentially activatable. Also, a method of retooling genes and gene pathways by the introduction of regulatory sequences, such as promoters, that are operable in an intended host, thus conferring operability to a novel gene pathway when it is introduced into an intended host. For example a novel man-made gene pathway, generated based on microbially-derived progenitor templates, that is operable in a plant cell. Furthermore, a method of generating novel host organisms having increased expression of desirable traits, recombinant genes, and gene products.

CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 4 OF 19 USPATFULL on STN

```
AN
       2005:62926 USPATFULL
ΤI
       Amplification of nucleic acids with electronic detection
IN
       Blackburn, Gary, Glendora, CA, UNITED STATES
       Irvine, Bruce D., Glendora, CA, UNITED STATES
       Kavvem, Jon Faiz, Pasadena, CA, UNITED STATES
       Sheldon, Edward Lewis, III, Arcadia, CA, UNITED STATES
       Terbrueggen, Robert H., Manhattan Beach, CA, UNITED STATES
PΙ
       US 20050053962
                         A1 20050310
                          A1 20041115 (10)
      US 2004-746904
```

ΑI Continuation of Ser. No. US 2000-621275, filed on 20 Jul 2000, GRANTED, RLI Pat. No. US 6686150 Continuation-in-part of Ser. No. US 1999-238351, filed on 27 Jan 1999, PENDING Continuation-in-part of Ser. No. US 1998-14304, filed on 27 Jan 1998, GRANTED, Pat. No. US 6063573 Continuation-in-part of Ser. No. US 1998-135183, filed on 17 Aug 1998, PENDING

DT Utility FS

L8

APPLICATION LREP DORSEY & WHITNEY LLP, INTELLECTUAL PROPERTY DEPARTMENT, 4 EMBARCADERO CENTER, SUITE 3400, SAN FRANCISCO, CA, 94111

CLMN Number of Claims: 25 ECI. Exemplary Claim: 1 DRWN 66 Drawing Page(s) IN. CNT 6587

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The invention relates to compositions and methods useful in the detection of nucleic acids using a variety of amplification techniques, including both signal amplification and target amplification. Detection proceeds through the use of an electron transfer moiety (ETM) that is associated with the nucleic acid, either directly or indirectly, to allow electronic detection of the ETM using an electrode.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

1.8 ANSWER 5 OF 19 USPATFULL on STN

AN

2004:101228 USPATFULL

ΤI Whole cell engineering by mutagenizing a substantial portion of a starting genome, combining mutations, and optionally repeating

IN Short, Jay M., Rancho Santa Fe, CA, UNITED STATES PΙ

US 20040077090 A1 20040422

US 2003-383798 A1 20030306 (10) AΙ

RLI Continuation of Ser. No. US 2000-677584, filed on 30 Sep 2000, ABANDONED Continuation-in-part of Ser. No. US 2000-594459, filed on 14 Jun 2000, GRANTED, Pat. No. US 6605449 Continuation-in-part of Ser. No. US 2000-522289, filed on 9 Mar 2000, GRANTED, Pat. No. US 6358709 Continuation-in-part of Ser. No. US 2000-498557, filed on 4 Feb 2000, PENDING Continuation-in-part of Ser. No. US 2000-495052, filed on 31 Jan 2000, GRANTED, Pat. No. US 6479258

PRAI US 1999-156815P 19990929 (60)

Utility DT

FS APPLICATION

LREP HALE AND DORR LLP, 300 PARK AVENUE, NEW YORK, NY, 10022

CLMN Number of Claims: 22 ECL Exemplary Claim: 1

DRWN 28 Drawing Page(s)

LN.CNT 37121

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB An invention comprising cellular transformation, directed evolution, and screening methods for creating novel transgenic organisms having desirable properties. Thus in one aspect, this invention relates to a method of generating a transgenic organism, such as a microbe or a plant, having a plurality of traits that are differentially activatable. Also, a method of retooling genes and gene pathways by the introduction of regulatory sequences, such as promoters, that are operable in an intended host, thus conferring operability to a novel gene pathway when it is introduced into an intended host. For example a novel man-made gene pathway, generated based on microbially-derived progenitor templates, that is operable in a plant cell. Furthermore, a method of generating novel host organisms having increased expression of desirable traits, recombinant genes, and gene products.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 6 OF 19 USPATFULL on STN

2004:27054 USPATFULL AN

Amplification of nucleic acids with electronic detection

ΤN Blackburn, Gary, Glendora, CA, United States Irvine, Bruce D., Glendora, CA, United States Kayyem, Jon Faiz, Pasadena, CA, United States Sheldon, III, Edward Lewis, Arcadia, CA, United States Terbrueggen, Robert H., Manhattan Beach, CA, United States

```
PA
       Clinical Micro Sensors, Inc., Pasadena, CA, United States (U.S.
       corporation)
                           B1 20040203
PΤ
       US 6686150
       US 2000-621275
                               20000720 (9)
AΙ
RLI
       Continuation-in-part of Ser. No. US 1999-238351, filed on 27 Jan 1999
       Continuation of Ser. No. US 1998-14304, filed on 27 Jan 1998, now
       patented, Pat. No. US 6063573 Continuation of Ser. No. US 1998-135183,
       filed on 17 Aug 1998
PRAI
       US 1999-144698P
                               19990720 (60)
       US 1998-84425P
                               19980506 (60)
       US 1998-84509P
                               19980506 (60)
       US 1998-28102P
                               19980316 (60)
       US 1998-73011P
                               19980129 (60)
DT
       Utility
FS
       GRANTED
EXNAM Primary Examiner: Marschel, Ardin H.
       Dorsey & Whitney LLP, Silva, Robin M., Kosslak, Renee M.
CLMN
       Number of Claims: 23
ECL
       Exemplary Claim: 1
DRWN
       104 Drawing Figure(s); 66 Drawing Page(s)
LN.CNT 7336
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AB
       The invention relates to compositions and methods useful in the
       detection of nucleic acids using a variety of amplification techniques.
       including both signal amplification and target amplification. Detection
       proceeds through the use of an electron transfer moiety (ETM) that is
       associated with the nucleic acid, either directly or indirectly, to
       allow electronic detection of the ETM using an electrode.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
     ANSWER 7 OF 19 USPATFULL on STN
L8
       2003:127016 USPATFULL
AN
ΤI
       ELECTRONIC DETECTION OF NUCLEIC ACIDS USING MONOLAYERS
       BAMDAD, CYNTHIA, SHARON, MA, UNITED STATES
ΤN
       YU, CHANGJUN, PASADENA, CA, UNITED STATES
ΡI
       US 20030087228
                          A1 20030508
ΑI
       US 1999-245105
                           A1 19990127 (9)
PRAI
       US 1998-84425P
                               19980506 (60)
       US 1998-84509P
                               19980506 (60)
       Utility
FS
       APPLICATION
LREP
       FLEHR HOHBACH TEST ALBRITTON & HERBERT, ROBIN M SILVA, SUITE 3400 FOUR
       EMBARCADERO CENTER, SAN FRANCISCO, CA, 941114187
CLMN
       Number of Claims: 38
ECL
       Exemplary Claim: 1
DRWN
      50 Drawing Page(s)
LN.CNT 4573
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       The present invention is directed to the electronic detection of nucleic
AB
       acids using self-assembled monolayers.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
     ANSWER 8 OF 19 USPATFULL on STN
```

Method and system for array signal generation and amplification

Gellibolian, Robert, Fremont, CA, UNITED STATES

A1 20010817 (9)

ΔNI

TT

TN

DT

AI

2003:51108 USPATFULL

US 2001-932728

Utility

US 20030036065 A1 20030220

```
APPLICATION
LREP
      AGILENT TECHNOLOGIES, INC., Intellectual Property Administration, Legal
       Department, DL429, P.O. Box 7599, Loveland, CO, 80537-0599
CLMN
      Number of Claims: 24
ECL
      Exemplary Claim: 1
DRWN 16 Drawing Page(s)
LN.CNT 885
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       A method and system for signal generation and signal amplification from
       an array containing bound, unlabeled target molecules. Following
       exposure of the array to a sample solution containing unlabaled target
       RNA molecules, blunt ends are generated on each probe/target
       double-stranded hybrid labeled primer oligonucleotide
       linker is then bound to the blunt ends. Next, in an iterative,
       inner process, additional layers of labeled oligonucleotide, linkers are
       added, shell-by-shell, to form a dendrimer-like molecular complex bound
       through the oligonucleotide linker to the probe/target hybrid.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
    ANSWER 9 OF 19 USPATFULL on STN
1.8
       2002:221316 USPATFULL
AN
TI
      Methods and products for analyzing polymers
       Chan, Eugene Y., Brookline, MA, UNITED STATES
IN
PΙ
       US 20020119455
                          A1 20020829
       US 2001-852968
                              20010510 (9)
AΤ
                          A1
RLT
       Division of Ser. No. US 1998-134411, filed on 13 Aug 1998, PATENTED
      WO 1998-US3024
PRAI
                               19980211
      US 1997-64687P
                               19971105 (60)
      US 1997-37921P
                              19970212 (60)
DТ
      Utility
FS
      APPLICATION
LREP
      Helen C. Lockhart, Esq., Wolf, Greenfield & Sacks, P.C., 600 Atlantic
      Avenue, Boston, MA, 02210
CLMN
      Number of Claims: 159
ECL
      Exemplary Claim: 1
DRWN
      10 Drawing Page(s)
LN.CNT 6864
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
      Methods and products for analyzing polymers are provided. The methods
       include methods for determining various other structural properties of
       the polymers.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
1.8
     ANSWER 10 OF 19 USPATFULL on STN
AN
       2002:12264 USPATFULL
ΤI
       AMPLIFICATION OF NUCLEIC ACIDS WITH ELECTRONIC DETECTION
       KAYYEM, JON FAIZ, PASADENA, CA, UNITED STATES
TN
       BAMDAD, CYNTHIA, SAN MARINO, CA, UNITED STATES
ΡI
       US 20020006643
                          A1 20020117
       US 7090804
                           B2 20060815
                          A1 19990127 (9)
AΙ
      US 1999-238351
       Continuation of Ser. No. US 1998-14304, filed on 27 Jan 1998, GRANTED,
RLI
       Pat. No. US 6063573 Continuation of Ser. No. US 1998-135183, filed on 17
       Aug 1998, PENDING
PRAT
      US 1998-84425P
                               19980506 (60)
       US 1998-84509P
                              19980506 (60)
       US 1996-28102P
                              19961009 (60)
      US 1998-73011P
                              19980129 (60)
      Utility
```

```
FS
       APPLICATION
LREP
       FLEHR HOHBACH TEST ALBRITTON & HERBERT, SUITE 3400, FOUR EMBARCADERO
       CENTER, SAN FRANCISCO, CA, 941114187
CLMN
       Number of Claims: 20
ECL
      Exemplary Claim: 1
DRWN 60 Drawing Page(s)
LN.CNT 5702
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       The invention relates to compositions and methods useful in the
       detection of nucleic acids using a variety of amplification techniques.
       including both signal amplification and target amplification. Detection
       proceeds through the use of an electron transfer moiety (ETM) that is
       associated with the nucleic acid, either directly or indirectly, to
       allow electronic detection of the ETM using an electrode.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
T.R
     ANSWER 11 OF 19 USPATFULL on STN
AN
       2002:50774 USPATFULL
       Methods and products for analyzing polymers
       Chan, Eugene Y., Brookline, MA, United States
       US Genomics, Woburn, MA, United States (U.S. corporation)
PA
                           B1 20020312
PΙ
       US 6355420
ΑI
       US 1998-134411
                               19980813 (9)
RLI
       Continuation of Ser. No. WO 1998-US3024, filed on 11 Feb 1998
       US 1997-37921P
                               19970212 (60)
PRAT
       US 1997-64687P
                               19971105 (60)
       Htility
FS
       GRANTED
EXNAM Primary Examiner: Jones, W. Gary; Assistant Examiner: Taylor, Janell E.
      Wolf, Greenfield & Sacks, P.C.
LREP
CLMN
      Number of Claims: 123
ECL
      Exemplary Claim: 1
DRWN
      15 Drawing Figure(s); 10 Drawing Page(s)
LN.CNT 6818
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       Methods and products for analyzing polymers are provided. The methods
       include methods for determining various other structural properties of
       the polymers.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
T.R
     ANSWER 12 OF 19 USPATFULL on STN
AN
       2001:67866 USPATFULL
TI
       Synthesis of fluorinated xanthene derivatives
       Klaubert, Dieter H., Sunnyvale, CA, United States
TN
       Gee, Kyle R., Eugene, OR, United States
PA
       Molecular Probes, Inc., Eugene, OR, United States (U.S. corporation)
                           B1 20010508
ΡI
       US 6229055
       US 2000-632251
AΙ
                               20000803 (9)
       Division of Ser. No. US 1996-631202, filed on 12 Apr 1996, now patented,
       Pat. No. US 6162931
       Utility
FS
       Granted
EXNAM Primary Examiner: Shippen, Michael L.
      Helfenstein, Allega T.
CLMN
      Number of Claims: 16
ECI.
       Exemplary Claim: 1
DRWN
     5 Drawing Figure(s); 5 Drawing Page(s)
LN.CNT 4318
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
```

AB Facile syntheses for fluorinated resorcinol and aminophenol derivatives are provided that yield isomer-free products in good yield. These novel methods use generally available precursors and standard laboratory reagents and equipment to reproducibly produce these synthetically useful reagents in relatively large quantities. The resulting fluorinated resorcinols and aniinophenols possess utility in the preparation of fluorinated fluorescein and rhodol dyes.

# CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 13 OF 19 USPATFULL on STN

2001:1856 USPATFULL

L8

 $\Delta M$ 

```
ΤI
       PNA probes for detection of Neisseria gonorrhoeae and Chlamydia
       trachomatis
TN
       Hyldig-Nielsen, Jens J.o slashed.rgen, Vanl.o slashed.se, Denmark
       Godskesen, Michael Anders, Vedb.ae butted.k, Denmark
       DAKO A/S, Glostrup, Denmark (non-U.S. corporation)
PA
PΙ
       US 6169169
                          B1 20010102
      US 1995-443930
ΑI
                               19950518 (8)
PRAI
      DK 1994-572
                               19940519
      Utility
      Granted
EXNAM Primary Examiner: Riley, Jezia
LREP
      Graham & James LLP
CLMN
      Number of Claims: 15
ECL
      Exemplary Claim: 1
DRWN
      No Drawings
LN.CNT 1440
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AB
       Specific peptide nucleic acid (PNA) probes for detecting a sexual
       transmitted disease caused by Neisseria gonorrhoeae or Chlamydia
       trachomatis comprising N-(2-aminoethyl)glycine units in amide linkage
       with the glycine nitrogen connected to naturally occurring nucleobases
       or non-naturally occurring nucleobases by a methylene carbonyl linker
       and said probes capable of hybridizing to 16S or 23S rRNA or rDNA of
      Neisseria gonorrhoeae or Chlamydia trachomatis are described. PNA is a
       very stable molecule with very high affinity for nucleic acid allowing a
       PNA probe to be shorter than conventional nucleic acid probes.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L8
     ANSWER 14 OF 19 USPATFULL on STN
AN
       2000:171155 USPATFULL
ΤI
       Fluorinated xanthene derivatives
IN
       Gee, Kyle R., Springfield, OR, United States
       Poot, Martin, Eugene, OR, United States
       Klaubert, Dieter H., Eugene, OR, United States
       Sun, Wei-Chuan, Eugene, OR, United States
       Haugland, Richard P., Eugene, OR, United States
       Mao, Fei, Eugene, OR, United States
       Molecular Probes, Inc., Eugene, OR, United States (U.S. corporation)
PA
PΙ
      US 6162931
                               20001219
      US 1996-631202
ΑI
                               19960412 (8)
DT
      Utility
FS
      Granted
EXNAM Primary Examiner: Riley, Jezia
LREP
      Skaugset, Anton E., Helfenstein, Allegra J.
CLMN
      Number of Claims: 107
ECI.
       Exemplary Claim: 1
DRWN 5 Drawing Figure(s); 5 Drawing Page(s)
LN.CNT 5371
```

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The family of dyes of the invention are fluoresceins and rhodols that AB are directly substituted on one or more aromatic carbons by fluorine. These fluorine-substituted fluorescent dyes possess greater photostability and have lower sensitivity to pH changes in the physiological range of 6-8 than do non-fluorinated dyes, exhibit less quenching when conjugated to a substance, and possess additional advantages. The dyes of the invention are useful as detectable tracers and for preparing conjugates of organic and inorganic substances.

#### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L8 ANSWER 15 OF 19 USPATFULL on STN

AN 2000:24765 USPATFULL

ΤТ Non-nucleotide linking reagents for nucleotide

probes

TN Arnold, Jr., Lyle J., Poway, CA, United States Reynolds, Mark A., Lafayette, CO, United States Bhatt, Ram S., San Diego, CA, United States

PA Gen-Probe Incorporated, San Diego, CA, United States (U.S. corporation) ΡI

US 6031091 20000229

ΑI US 1997-908535 19970807 (8)

Continuation-in-part of Ser. No. US 1995-485629, filed on 7 Jun 1995, RLI now patented, Pat. No. US 5696251 which is a division of Ser. No. US 1994-182666, filed on 14 Jan 1994, now patented, Pat. No. US 5585481 which is a continuation of Ser. No. US 1989-319422, filed on 6 Mar 1989, now abandoned which is a continuation-in-part of Ser. No. US 1987-99050, filed on 21 Sep 1987, now abandoned

DT Utility

FS Granted

EXNAM Primary Examiner: Wilson, James O.

LREP Lyon & Lyon LLP

CLMN Number of Claims: 32

ECL Exemplary Claim: 1

DRWN 20 Drawing Figure(s); 20 Drawing Page(s)

LN.CNT 1547

AB

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A versatile reagent with a non-nucleotide monomeric unit having a ligand, and first and second coupling groups which are linked to the non-nucleotide monomeric unit. The ligand can be either a chemical moiety, such as a label or intercalator, or a linking arm which can be linked to such a moiety. Such reagent permits preparation of versatile nucleotide/non-nucleotide polymers, having any desired sequence of nucleotide and nonnucleotide monomeric units, each of the latter of which bear a desired ligand. These polymers can for example, be used as probes which can exhibit enhanced sensitivity and/or which are capable of detecting a genus of nucleotides each species of which has a common target nucleotide sequence of interest bridged by different sequences not of interest.

#### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 16 OF 19 USPATFULL on STN

1999:146263 USPATFULL AN

Detection of Ribosomal RNA using PNA probes

TN Hyldig-Nielsen, Jens J.o slashed.rgen, Vanl.o slashed.se, Denmark

Godskesen, Michael Anders, Vedb.ae butted.k, Denmark PΆ Dako A/S, Glostrup, Denmark (non-U.S. corporation)

PT US 5985563 19991116

US 1997-869454 AΤ 19970605 (8)

```
Division of Ser. No. US 1995-443930, filed on 18 May 1995
```

PRAT DK 1994-572 19940519

Utility DT FS

Granted

EXNAM Primary Examiner: Marschel, Ardin H.: Assistant Examiner: Riley, Jezia LREP Graham & James LLP

CLMN Number of Claims: 38

ECL Exemplary Claim: 1 DRWN No Drawings

LN.CNT 1564

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

PNA probes for detection of Neisseria gonorrhoeae and Chlamydia trachomatis.

Specific peptide nucleic acid (PNA) probes for detecting a sexual transmitted disease caused by Neisseria gonorrhoeae or Chlamydia trachomatis comprising N-(2-aminoethyl)glycine units in amide linkage with the glycine nitrogen connected to naturally occurring nucleobases or non-naturally occurring nucleobases by a methylene carbonyl linker and said probes capable of hybridizing to 16S or 23S rRNA or rDNA of Neisseria gonorrhoeae or Chlamydia trachomatis are described.

PNA is a very stable molecule with very high affinity for nucleic acid allowing a PNA probe to be shorter than conventional nucleic acid probes.

#### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 17 OF 19 USPATFULL on STN

AN 97:115408 USPATFULL

TI Non-nucleotide linking reagents for nucleotide

probes

Arnold, Jr., Lyle J., San Diego, CA, United States Reynolds, Mark A., San Diego, CA, United States Bhatt, Ram S., San Diego, CA, United States

PA Gen-Probe Incorporated, San Diego, CA, United States (U.S. corporation) ΡI US 5696251 19971209

ΑI US 1995-485629 19950607 (8)

RLI

Continuation of Ser. No. US 1994-182666, filed on 14 Jan 1994, now patented, Pat. No. US 5585481 which is a continuation of Ser. No. US 1989-319422, filed on 6 Mar 1989, now abandoned which is a continuation-in-part of Ser. No. US 1987-99050, filed on 21 Sep 1987, now abandoned

PRAI

PT 1988-88550 19880920

DT Utility FS Granted

EXNAM Primary Examiner: Wilson, James O.

LREP Lyon & Lyon LLP

Number of Claims: 49 CLMN

ECL Exemplary Claim: 1

DRWN 20 Drawing Figure(s); 20 Drawing Page(s)

LN.CNT 1601

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A versatile reagent with a non-nucleotide monomeric unit

having a ligand, and first and second coupling groups which are linked to the non-nucleotide monomeric unit. The ligand can be either a chemical moiety, such as a label or intercalator, or a linking arm which can be linked to such a moiety. Such reagent permits preparation of versatile nucleotide/non-nucleotide polymers,

having any desired sequence of nucleotide and non-

nucleotide monomeric units, each of the latter of which bear a

desired ligand. These polymers can for example, be used as probes which can exhibit enhanced sensitivity and/or which are capable of detecting a genus of nucleotides each species of which has a common target nucleotide sequence of interest bridged by different sequences not of interest.

#### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

- ANSWER 18 OF 19 USPATFULL on STN L8
- AN 97:71177 USPATFULL
- ΤI Methods for making nucleotide polymers using novel linking
- reagents IN
- Arnold, Jr., Lyle J., San Diego, CA, United States Reynolds, Mark A., San Diego, CA, United States
- Bhatt, Ram S., San Diego, CA, United States
- Gen-Probe Incorporated, San Diego, CA, United States (U.S. corporation) PA
- PТ US 5656744 19970812 19950607 (8)
- US 1995-490109 ΑI
- RLI Division of Ser. No. US 1994-182666, filed on 14 Jan 1994, now patented, Pat. No. US 5585481 which is a continuation of Ser. No. US 1989-319422, filed on 6 Mar 1989, now abandoned which is a continuation-in-part of
- Ser. No. US 1987-99050, filed on 21 Sep 1987, now abandoned 19880920
- PT 1988-88550 PRAI
- DT Utility
- Granted EXNAM Primary Examiner: Wilson, James O.
- Lyon & Lyon LLP
- CLMN Number of Claims: 51
- ECL Exemplary Claim: 1
- 20 Drawing Figure(s); 20 Drawing Page(s)
- LN.CNT 1717
- CAS INDEXING IS AVAILABLE FOR THIS PATENT.
- A versatile reagent with a non-nucleotide monomeric unit
  - having a ligand, and first and second coupling groups which are linked to the non-nucleotide monomeric unit. The ligand can be either a chemical moiety, such as a label or intercalator, or a linking arm which can be linked to such a moiety. Such reagent permits preparation
  - of versatile nucleotide/non-nucleotide polymers, having any desired sequence of nucleotide and non-
  - nucleotide monomeric units, each of the latter of which bear a
  - desired ligand. These polymers can for example, be used as probes which can exhibit enhanced sensitivity and/or which are capable of detecting a genus of nucleotides each species of which has a common target
  - nucleotide sequence of interest bridged by different sequences not of interest.

#### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

- ANSWER 19 OF 19 USPATFULL on STN T.R
- AN 96:116488 USPATFULL
- ΤI Linking reagents for nucleotide probes
- IN Arnold, Jr., Lyle J., San Diego, CA, United States Reynolds, Mark A., San Diego, CA, United States
- Bhatt, Ram S., San Diego, CA, United States PA Gen-Probe Incorporated, San Diego, CA, United States (U.S. corporation)
- PI US 5585481 19961217
- AΤ US 1994-182666 19940114 (8)
- RLT Continuation of Ser. No. US 1989-319422, filed on 6 Mar 1989, now abandoned which is a continuation-in-part of Ser. No. US 1987-99050, filed on 21 Sep 1987, now abandoned
- PRAI PT 1988-88550 19880920

```
DT
      Utility
FS
       Granted
EXNAM Primary Examiner: Wilson, James O.
LREP Lvon & Lvon
CLMN Number of Claims: 50
ECL
      Exemplary Claim: 1
DRWN
       20 Drawing Figure(s); 20 Drawing Page(s)
LN.CNT 1715
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       A versatile reagent with a non-nucleotide monomeric unit
       having a ligand, and first and second coupling groups which are linked
       to the non-nucleotide monomeric unit. The ligand can be either
       a chemical moiety, such as a label or intercalator, or a linking arm
       which can be linked to such a moiety. Such reagent permits preparation
       of versatile nucleotide/non-nucleotide polymers,
       having any desired sequence of nucleotide and non-
       nucleotide monomeric units, each of the latter of which bear a
       desired ligand. These polymers can for example, be used as probes which
       can exhibit enhanced sensitivity and/or which are capable of detecting a
       genus of nucleotides each species of which has a common target
       nucleotide sequence of interest bridged by different sequences
       not of interest.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
=> d his
     (FILE 'HOME' ENTERED AT 11:49:49 ON 04 JUL 2010)
     FILE 'BIOSIS, MEDLINE, CAPLUS, WPIDS, USPATFULL' ENTERED AT 11:50:31 ON
     04 JUL 2010
           4804 S (NUCLEOTIDE OR NUCLEOSIDE) AND BASE (4A) (LABEL? OR MARKER OR
L2
            531 S L1 AND LINKER (4A) (LABEL? OR MARKER OR DYE)
L3
             27 S L2 AND LINKER (6A) POLYMER
L4
             0 S L3 AND WATER SOLUBLE POLYMER
L5
             8 S L3 AND LINKER (4A) 50
L6
             8 DUP REM L5 (0 DUPLICATES REMOVED)
L7
             19 S L3 NOT L6
1.8
             19 DUP REM L7 (0 DUPLICATES REMOVED)
=> s 12 and polymer (5a) water soluble
L9
             9 L2 AND POLYMER (5A) WATER SOLUBLE
=> dup rem 19
PROCESSING COMPLETED FOR L9
              9 DUP REM L9 (0 DUPLICATES REMOVED)
=> d 110 bib abs 1-9
L10 ANSWER 1 OF 9 USPATFULL on STN
       2004:94757 USPATFULL
AN
       Methods of labelling polynucleotides with dibenzorhodamine dves
ΤI
IN
       Benson, Scott C., Alameda, CA, UNITED STATES
       Lam, Joe Y.L., Castro Valley, CA, UNITED STATES
       Menchen, Steven Micheal, Fremont, CA, UNITED STATES
PA
       Applera Corporation, Foster City, CA, UNITED STATES, 94404 (U.S.
       corporation)
                          A1 20040415
PΤ
       US 20040072209
       US 6919445
                          B2 20050719
AΤ
       US 2003-441950
                          A1 20030520 (10)
```

- RLT Continuation of Ser. No. US 2001-969430, filed on 2 Oct 2001, GRANTED, Pat. No. US 6566071 Division of Ser. No. US 2001-784943, filed on 14 Feb 2001, GRANTED, Pat. No. US 6326153 Continuation of Ser. No. US 2000-556040, filed on 20 Apr 2000, GRANTED, Pat. No. US 6221606 Division of Ser. No. US 1998-199402, filed on 24 Nov 1998, GRANTED, Pat. No. US 6111116 Division of Ser. No. US 1997-978775, filed on 25 Nov 1997. GRANTED, Pat. No. US 5936087
- Utility
- FS APPLICATION LREP MILA KASAN, PATENT DEPT., APPLIED BIOSYSTEMS, 850 LINCOLN CENTRE DRIVE, FOSTER CITY, CA. 94404
- Number of Claims: 106 CLMN ECL Exemplary Claim: 1
- DRWN 7 Drawing Page(s)
- LN.CNT 1704
- CAS INDEXING IS AVAILABLE FOR THIS PATENT.
- Dibenzorhodamine compounds having the structure ##STR1##

are disclosed, including nitrogen- and aryl-substituted forms thereof. In addition, two intermediates useful for synthesizing such compounds are disclosed, a first intermediate having the structure

including nitrogen- and arvl-substituted forms thereof, and a second intermediate having the structure ##STR3##

including nitrogen- and aryl-substituted forms thereof, wherein substituents at positions C-14 to C18 taken separately are selected from the group consisting of hydrogen, chlorine, fluorine, lower alkyl, carboxylic acid, sulfonic acid, --CH.sub.20H, alkoxy, phenoxy, linking group, and substituted forms thereof. The invention further includes energy transfer dyes comprising the dibenzorhodamine compounds, nucleosides labeled with the dibenzorhodamine compounds, and nucleic acid analysis methods employing the dibenzorhodamine compounds.

## CAS INDEXING IS AVAILABLE FOR THIS PATENT.

- L10 ANSWER 2 OF 9 USPATFULL on STN
- AN 2002:60927 USPATFULL
- ΤI Methods of labelling polynucleotides with dibenzorhodamine dyes
- TN Benson, Scott C., Alameda, CA, UNITED STATES Lam, Joe Y.L., Castro Valley, CA, UNITED STATES
- Menchen, Steven Michael, Fremont, CA, UNITED STATES
- PA The Perkin-Elmer Corporation, Foster City, CA, UNITED STATES, 94404 (U.S. corporation)
- PΙ US 20020034761 A1 20020321 US 6566071 B2 20030520
- US 2001-969430 A1 20011002 (9) AΤ
- Division of Ser. No. US 2001-784943, filed on 14 Feb 2001, PENDING RLI Continuation of Ser. No. US 2000-556040, filed on 20 Apr 2000, GRANTED, Pat. No. US 6221606 Division of Ser. No. US 1998-199402, filed on 24 Nov 1998, GRANTED, Pat. No. US 6111116 Division of Ser. No. US 1997-978775, filed on 25 Nov 1997, GRANTED, Pat. No. US 5936087
- Utility
- FS APPLICATION
- PATTI SELAN, PATENT ADMINISTRATOR, APPLIED BIOSYSTEMS, 850 LINCOLN LREP CENTRE DRIVE, FOSTER CITY, CA, 94404
- CLMN Number of Claims: 106
- ECL Exemplary Claim: 1 DRWN 7 Drawing Page(s)
- LN.CNT 1703

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB

are disclosed, including nitrogen- and aryl-substituted forms thereof. In addition, two intermediates useful for synthesizing such compounds are disclosed, a first intermediate having the structure #\$STR2#

including nitrogen- and aryl-substituted forms thereof, and a second intermediate having the structure ##STR3##

including nitrogen— and aryl-substituted forms thereof, where in substituents at positions C-14 to C18 taken separately are selected from the group consisting of hydrogen, chlorine, fluorine, lower alkyl, carboxylic acid, sulfonic acid, --CH.sub.20H, alkoxy, phenoxy, linking group, and substituted forms thereof. The invention further includes energy transfer dyes comprising the dibenzorhodamine compounds, nucleosides labeled with the dibenzorhodamine compounds, and nucleic acid analysis methods employing the dibenzorhodamine compounds.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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L10 ANSWER 3 OF 9 USPATFULL on STN
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AN 2001:123631 USPATFULL

TI Polynucleotides labeled with dibenzorhodamine dyes

IN Benson, Scott C., Alameda, CA, United States Lam, Joe Y.L., Castro Valley, CA, United States

Menchen, Steven Michael, Fremont, CA, United States
PA The Perkin-Elmer Corporation (U.S. corporation)

PI US 20010011139 A1 20010802

US 20010011139 A1 20010802 US 6326153 B2 20011204

AI US 2001-784943 A1 20010214 (9)

RLI Continuation of Ser. No. US 2000-556040, filed on 20 Apr 2000, GRANTED, Pat. No. US 6221606 Division of Ser. No. US 1998-199402, filed on 24 Nov 1998, GRANTED, Pat. No. US 6111116 Division of Ser. No. US 1997-978775,

filed on 25 Nov 1997, GRANTED, Pat. No. US 5936087

DT Utility

FS APPLICATION

LREP PATTI SELAN, PATENT ADMINISTRATOR, APPLIED BIOSYSTEMS, 850 LINCOLN CENTRE DRIVE, FOSTER CITY, CA, 94404

CLMN Number of Claims: 106

ECL Exemplary Claim: 1 DRWN 7 Drawing Page(s)

LN.CNT 1687

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Dibenzorhodamine compounds having the structure ##STR1##

are disclosed, including nitogen— and aryl-substituted forms thereof. In addition, two intermediates useful for synthesizing such compounds are disclosed, a first intermediate having the structure  $$\#\$5\mbox{TR}2\#\$$$ 

including nitrogen- and aryl-substituted forms thereof, and a second intermediate having the structure ##STR3##

including nitrogen- and aryl-substituted forms thereof, wherein substituents at positions C-14 to C18 taken separately are selected from the group consisting of hydrogen, chlorine, fluorine, lower alkyl, carboxylic acid, sulfonic acid, --CH.sub.20H, alkoxy, phenoxy, linking group, and substituted forms thereof. The invention further includes energy transfer dyes comprising the dibenzorhodamine compounds, nucleosides labeled with the dibenzorhodamine compounds, and nucleic acid analysis methods employing the dibenzorhodamine compounds.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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L10 ANSWER 4 OF 9 USPATFULL on STN
```

- 2001:59629 USPATFULL AN
- TI Dibenzorhodamine dves
- IN Benson, Scott C., Foster City, CA, United States
  - Lam, Joe Y. L., Foster City, CA, United States
- Menchen, Steven Michael, Foster City, CA, United States The Perkin-Elmer Corporation, Foster City, CA, United States (U.S.
- corporation)
- PΙ US 6221606 B1 20010424
- AΙ US 2000-556040 20000420 (9)
- RLI Division of Ser. No. US 1998-199402, filed on 24 Nov 1998, now patented, Pat. No. US 6111116 Division of Ser. No. US 1997-978775, filed on 25 Nov 1997, now patented, Pat. No. US 5936087
- DТ Utility FS Granted
- EXNAM Primary Examiner: Davis, Zinna Northington
- Andrus, Alex, Grossman, Paul D.
- CLMN Number of Claims: 52
- ECL Exemplary Claim: 1
- DRWN 7 Drawing Figure(s); 7 Drawing Page(s)
- LN.CNT 1526
- CAS INDEXING IS AVAILABLE FOR THIS PATENT.
- Dibenzorhodamine compounds having the structure ##STR1##

are disclosed, including nitogen- and aryl-substituted forms thereof. In addition, two intermediates useful for synthesizing such compounds are disclosed, a first intermediate having the structure ##STR2##

including nitrogen- and aryl-substituted forns thereof, and a second intermediate having the structure ##STR3##

including nitrogen- and aryl-substituted forms thereof, wherein substituents at positions C-14 to C18 taken separately are selected from the group consisting of hydrogen, chlorine, fluorine, lower alky, carboxylic acid, sulfonic acid, --CH.sub.2 OH, alkoxy, phenoxy, linking group, and substituted forms thereof. The invention further includes energy transfer dyes comprising the dibenzorhodamine compounds, nucleosides labeled with the dibenzorhodamine compounds, and nucleic acid analysis methods employing the dibenzorhodamine compounds.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

- L10 ANSWER 5 OF 9 USPATFULL on STN
- AN 2001:1856 USPATFULL
- ΤТ PNA probes for detection of Neisseria gonorrhoeae and Chlamydia trachomatis
- TN Hyldig-Nielsen, Jens J.o slashed.rgen, Vanl.o slashed.se, Denmark Godskesen, Michael Anders, Vedb.ae butted.k, Denmark
  - DAKO A/S, Glostrup, Denmark (non-U.S. corporation)
- PA PΙ US 6169169 B1 20010102
- US 1995-443930 ΑI 19950518 (8) 19940519
- DK 1994-572 PRAI DT Utility
- FS Granted EXNAM Primary Examiner: Riley, Jezia
- LREP Graham & James LLP CLMN Number of Claims: 15
- ECL Exemplary Claim: 1
- DRWN No Drawings

LN.CNT 1440

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Specific peptide nucleic acid (PNA) probes for detecting a sexual transmitted disease caused by Neisseria gonorrhoeae or Chlamydia trachomatis comprising N-(2-aminoethyl)glycine units in amide linkage with the glycine nitrogen connected to naturally occurring nucleobases or non-naturally occurring nucleobases by a methylene carbonyl linker and said probes capable of hybridizing to 16S or 23S rRNA or rDNA of Neisseria gonorrhoeae or Chlamydia trachomatis are described. PNA is a very stable molecule with very high affinity for nucleic acid allowing a PNA probe to be shorter than conventional nucleic acid probes.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L10 ANSWER 6 OF 9 USPATFULL on STN

AN 2000:114142 USPATFULL

тт Dibenzorhodamine dves

IN Benson, Scott C., Foster City, CA, United States Lam, Joe Y. L., Foster City, CA, United States

Menchen, Steven Michael, Foster City, CA, United States

The Perkin-Elmer Corporation, Foster City, CA, United States (U.S. PA corporation) 20000829

US 6111116 PT

ΑI US 1998-199402

19981124 (9) RLI Division of Ser. No. US 1997-978775, filed on 25 Nov 1997, now patented, Pat. No. US 5936087

Utility

Granted

EXNAM Primary Examiner: Davis, Zinna Northington

LREP Grossman, Paul D., Andrus, Alex

CLMN Number of Claims: 44

ECL Exemplary Claim: 1

DRWN 7 Drawing Figure(s); 7 Drawing Page(s) LN.CNT 1501

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Dibenzorhodamine compounds having the structure ##STR1## are disclosed, including nitogen- and aryl-substituted forms thereof. In addition, two intermediates useful for synthesizing such compounds are disclosed, a first intermediate having the structure ##STR2## including nitrogen- and aryl-substituted forms thereof, and a second intermediate having the structure ##STR3## including nitrogen- and aryl-substituted forms thereof, wherein substituents at positions C-14 to C18 taken separately are selected from the group consisting of hydrogen, chlorine, fluorine, lower alkyl, carboxylic acid, sulfonic acid, --CH.sub.2 OH, alkoxy, phenoxy, linking group, and substituted forms thereof. The invention further includes energy transfer dyes comprising the dibenzorhodamine compounds, nucleosides labeled with the dibenzorhodarnine compounds, and nucleic acid analysis methods employing the dibenzorhodamine compounds.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L10 ANSWER 7 OF 9 USPATFULL on STN

2000:47376 USPATFULL AN

TI Dibenzorhodamine dves

IN Benson, Scott Conrad, Oakland, CA, United States Lam, Joe Y. L., Castro Valley, CA, United States Upadhya, Krishna Gajanan, Union City, CA, United States Radel, Peggy Ann, Berkeley, CA, United States Zhen, Weiguo, Foster City, CA, United States Menchen, Steven Michael, Fremont, CA, United States

PΆ The Perkin-Elmer Corporation, Foster City, CA, United States (U.S.

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corporation)
PΤ
      US 6051719
                               20000418
      US 1998-193374
                               19981117 (9)
AΤ
      Continuation-in-part of Ser. No. US 1997-978775, filed on 25 Nov 1997,
RLI
       now patented, Pat. No. US 5936087
      Utility
FS
      Granted
EXNAM Primary Examiner: Davis, Zinna Northington
LREP Andrus, Alex, Grossman, Paul D.
CLMN
      Number of Claims: 51
ECL
      Exemplary Claim: 1
DRWN
     15 Drawing Figure(s); 15 Drawing Page(s)
```

LN.CNT 1852

CAS INDEXING IS AVAILABLE FOR THIS PATENT. AB

Dibenzorhodamine compounds having the structure ##STR1## are disclosed, including nitogen- and aryl-substituted forms thereof. In addition, intermediates useful for synthesizing such compounds are disclosed, such intermediates having the structure ##STR2## In Formula I, R.sub.1 is H or ##STR3## wherein Y is H, lower alkyl, lower alkene, lower alkyne, aromatic, phenyl, polycyclic aromatic, heterocycle, water-solubilizing group, or linking group, including substituted forms thereof. When R.sub.1 is H, the C-12-bonded nitrogen and the C-12 and C-13 carbons form a first ring structure having from 4 to 7 members, and/or the C-12-bonded nitrogen and the C-11 and C-12 carbons form a second ring structure having from 5 to 7 members. The compounds of Formula I further include aryl- and nitrogen-substituted forms thereof. The invention further includes energy transfer dyes comprising the dibenzorhodamine compounds, nucleosides labeled with the dibenzorhodamine compounds, and nucleic acid analysis methods employing the dibenzorhodamine compounds.

### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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L10 ANSWER 8 OF 9 USPATFULL on STN
       1999:146263 USPATFULL
AN
ΤI
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Detection of Ribosomal RNA using PNA probes

TN Hyldig-Nielsen, Jens J.o slashed.rgen, Vanl.o slashed.se, Denmark Godskesen, Michael Anders, Vedb.ae butted.k, Denmark

PA Dako A/S, Glostrup, Denmark (non-U.S. corporation) PΙ US 5985563 19991116

US 1997-869454 19970605 (8)

ΑI RLI Division of Ser. No. US 1995-443930, filed on 18 May 1995

PRAI DK 1994-572 19940519

DT Utility

FS Granted EXNAM Primary Examiner: Marschel, Ardin H.; Assistant Examiner: Riley, Jezia LREP Graham & James LLP

CLMN Number of Claims: 38 ECL Exemplary Claim: 1

DRWN No Drawings LN.CNT 1564

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB PNA probes for detection of Neisseria gonorrhoeae and Chlamydia trachomatis.

Specific peptide nucleic acid (PNA) probes for detecting a sexual transmitted disease caused by Neisseria gonorrhoeae or Chlamydia trachomatis comprising N-(2-aminoethyl)glycine units in amide linkage with the glycine nitrogen connected to naturally occurring nucleobases or non-naturally occurring nucleobases by a methylene carbonyl linker and said probes capable of hybridizing to 16S or 23S rRNA or rDNA of Neisseria gonorrhoeae or Chlamydia trachomatis are described.

PNA is a very stable molecule with very high affinity for nucleic acid allowing a PNA probe to be shorter than conventional nucleic acid probes.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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L10 ANSWER 9 OF 9 USPATFULL on STN
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AN 1999:92800 USPATFULL

TI Dibenzorhodamine dves

IN Benson, Scott C., Foster City, CA, United States Lam, Joe Y. L., Foster City, CA, United States Menchen, Steven Michael, Foster City, CA, United States

PA The Perkin-Elmer Corporation, Foster City, CA, United States (U.S.

corporation)

US 5936087 19990810 US 1997-978775 19971125 (8)

AI US 1997-DT Utility

FS Granted

EXNAM Primary Examiner: Davis, Zinna Northington

LREP Grossman, Paul D.

CLMN Number of Claims: 40 ECL Exemplary Claim: 1

DRWN 7 Drawing Figure(s); 7 Drawing Page(s)

LN.CNT 1459

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Dibenzorhodamine compounds having the structure ##STR1## are disclosed, including nitogen- and aryl-substituted forms thereof. In addition, two intermediates useful for synthesizing such compounds are disclosed, a first intermediate having the structure ##STR2## including nitrogen- and aryl-substituted forms thereof, and a second intermediate having the structure ##STR3## including nitrogen- and aryl-substituted forms thereof, entering substituents at positions C-14 to C18 taken separately are selected from the group consisting of hydrogen, chlorine, fluorine, lower alkyl, carboxylic acid, sulfonic acid, --CH.sub.2 OH, alkoxy, phenoxy, linking group, and substituted forms thereof. The invention further includes energy transfer dyes comprising the dibenzorhodamine compounds, and nucleosides labeled with the dibenzorhodamine compounds, and nucleic acid analysis methods employing the dibenzorhodamine compounds.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

=> d 110 9 kwic

L10 ANSWER 9 OF 9 USPATFULL on STN

SUMM In a fourth aspect, the invention includes energy transfer dye compounds comprising a donor dye, an acceptor dye, and a linker linking the donor and acceptor dyes. The donor dye is capable of absorbing light at a first wavelength and emitting.

SUMM In a fifth aspect, the present invention includes labeled

nucleoside/tides having the structure SUMM wherein NUC is a nucleoside/tide or nucleoside/tide

analog and D is a dibenzorhodamine dye compound having the structure set forth above. According to the invention, NUC and. . . .

DEID "Nucleoside" refers to a compound consisting of a purine, deazapurine, or pyrimidine nucleoside base, e.g., adenine, guanine, cytosine, uracil, thymine, deazaadenine, deazaquanosine, and the like, linked to a pentose at the 1' position. When the nucleoside base is purine or "Jedeazapurine, the sugar moiety is attached at the 9-position of the purine or deazapurine, and when the

nucleoside base is pyrimidine, the sugar moiety is attached at the 1-position of the pyrimidine, e.g., Kornberg and Baker, DNA Replication, 2nd Ed. (Freeman, San Francisco, 1992). The term " nucleotide" as used herein refers to a phosphate ester of a nucleoside, e.g., triphosphate esters, wherein the most common site of esterification is the hydroxyl group attached to the C-5 position of the pentose. The term "nucleoside/tide" as used herein refers to a set of compounds including both nucleosides and nucleotides. "Analogs" in reference to nucleosides/tides include synthetic analogs having modified base moieties, modified sugar moieties and/or modified phosphate moieties, e.g. described generally by Scheit, Nucleotide Analogs (John Wiley, New York, 1980). Phosphate analogs comprise analogs of phosphate wherein the phosphorous atom is in the +5. . . alkoxy, e.g., methoxy, ethoxy, allyloxy, isopropoxy, butoxy, isobutoxy and phenoxy, amino or alkylamino, fluoro, chloro and bromo. The term "labeled nucleoside/tide" refers to nucleosides/tides which are covalently attached to the dye compounds of

- Formula I through a linkage.

  DEID "Polynucleotide" or "oligonucleotide" means polymers of natural nucleotide monomers or analogs thereof, including double and single stranded deoxyribonucleotides, ribonucleotides, α-anomeric forms thereof, and the like. Usually the nucleoside monomers are linked by phosphodiester linkages, where as used herein, the term "phosphodiester linkage" refers to phosphodiester bonds or bonds.

  DEID . of absorbing the excitation energy emitted by the donor dye and
- DEID

  . . or absorbing the exectation energy emitted by the donor dye an
  fluorescing at a second wavelength in response, and a linker
  which attaches the donor dye to the acceptor dye,
  the linker being effective to facilitate efficient energy
  transfer between the donor and acceptor dyes. A through discussion of
- the structure, synthesis.

  DETD . . a fused ring structure which is attached to the carbonyl carbon, and R.sub.28 includes a functional group which attaches the linker to the acceptor dve.
- DETD In another preferred embodiment of the energy-transfer-dye aspect of the present invention, the linker attaches to the dibenzorhodamine dye component of the energy transfer dye at the C-1 or 13 positions, or, alternatively, where the C-7 substituent is phenyl. . .
- DETD A. Nucleoside/tide Reagents
- DETD A preferred class of labeled reagents comprise nucleoside
  /tides that incorporate the dibenzorhodamine dyes of the invention. Such
  nucleoside/tide reagents are particularly useful in the context
  of labeling polynucleotides formed by enzymatic synthesis, e.g.,
  nucleotide triphosphates used in the context of PCR
  amplification, Sanger-type polynucleotide sequencing, and
  nick-translation reactions.
- DETD Generally, the structure of the labeled nucleoside/tide reagent is
- DETD where NUC is a nucleoside/tide or nucleoside/tide analog and D is a dibenzorhodamine dye compound of Formula II.
- DETD The linkage linking the nucleoside/tide and the dye may be attached to the dye at any one of substituent positions C-1 to C-18 or at. . . nitrogen. Preferably, the dye includes a phenyl or
  - substituted phenyl substituent at the C-7 position and is attached to the nucleoside/tide at one of the C-15 or C-16 substituent positions, the other position being a hydrogen atom.
  - Nucleoside labeling can be accomplished using any one of a large number of known nucleoside/tide labeling techniques employing known linkages, linking groups, and associated complementary functionalities. Generally, the linkage linking the dye and nucleoside should (i) not interfere with oligonucleotide-target hybridization, (ii) be compatible with relevant enzymes, e.g.,

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polymerases, ligases, and the like, and (iii) not adversely affect the fluorescence properties of the dye. Exemplary base labeling procedures suitable for use in connection with the present invention include the following: Gibson et al, Nucleic Acids Research, 15:6455-6467.
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- DETD Preferably, the linkages are acetylenic amido or alkenic amido linkages, the linkage between the dye and the nucleoside/tide base being formed by reacting an activated N-hydroxysuccinimide (NHS) ester of the dye with an alkynylamino- or alkenylamino-derivatized
- base of a nucleoside/tide. More preferably, the resulting linkage is 3-(carboxy)amino-1-propyn-1-yl having the structure ##STR14## . . . for several hours, or until thin layer chromatography indicates consumption of the halodideoxynucleoside. When an unprotected
- consumption of the halodideoxynucleoside. When an unprotected alkynylamine is used, the alkynylamino-nucleoside can be isolated by concentrating the reaction mixture and chromatographing on silica gel using an eluting solvent which contains ammonium. . DETD Particularly preferred nucleosides/tides of the present invention are shown below in Formula IV wherein #\$STR18## B is a nucleoside /tide base, e.g., uracil, cytosine, deazaadenine, or deazaquanosine; W.sub.1 and W.sub.2 taken separately are OH or a group capable of
- DETD . . method, and the like, e.g., Gait, Oligonucleotide Synthesis, IRL Press (1990). Labels may be introduced during enzymatic synthesis utilizing labeled nucleotide triphosphate monomers as described above, or introduced during chemical synthesis using labeled non-nucleotide or nucleotide phosphoramidities as

blocking.

- described above, or may be introduced subsequent to synthesis. DETD . . . describes the steps of a typical polynucleotide synthesis cycle using the phosphoramidite method. First, a solid support including a protected nucleotide monomer is treated with acid, e.g., trichloroacetic acid, to remove a 5'-hydroxyl protecting group, freeing the hydroxyl for a subsequent coupling reaction. An activated intermediate is then formed by simultaneously adding a protected phosphoramidite nucleoside monomer and a weak acid, e.g., tetrazole, to the reaction. The weak acid protonates the nitrogen of the phosphoramidite forming a reactive intermediate. Nucleoside addition is complete within 30 s. Next, a capping step is performed which terminates any polynucleotide chains that did not undergo nucleoside addition. Capping is preferably done with acetic anhydride and 1-methylimidazole. The internucleotide linkage is then converted from the phosphite to.
- DETD Any of the phosphoramidite nucleoside monomers may be dye-labeled phosphoramidites as described above. If the 5'-terminal position of the nucleotide is labeled, a labeled non-nucleotidic phosphoramidite of the invention may be used during the final condensation step. If an internal.
- DETD . . . In the AmpFLPs technique, the polynucleotides may be labeled by using a labeled polynucleotide PCR primer, or by utilizing labeled nucleotide triphosphates in the PCR.
- DETD In another such fragment analysis method known as nick translation, a reaction is used to replace unlabeled nucleoside triphosphates in a double-stranded DNA molecule with labeled ones. Free 3'-hydroxyl groups are created within the unlabeled DNA by "nicks" caused by deoxyribonuclease I (DNAase I) treatment. DNA polymerase I then catalyzes the addition of a labeled nucleotide to the 3'-hydroxyl terminus of the nick. At the same time, the 5' to 3'-exonuclease activity of this enzyme eliminates the nucleotide unit from the 5'-phosphoryl terminus of the nick. A new nucleotide with a free 3'-DH group is incorporated at the position of the original excised nucleotide, and the nick is shifted along by one nucleotide unit in the 3' direction. This

```
3' shift will result in the sequential addition of new labeled
       nucleotides to the. . .
DETD
       . . . site based on where an oligonucleotide primer anneals to the
       template. The synthesis reaction is terminated by incorporation of a
       nucleotide analog that will not support continued DNA
       elongation. Exemplary chain-terminating nucleotide analogs
       include the 2',3'-dideoxynucleoside 5'-triphosphates (ddNTPs) which lack
       the 3'-OH group necessary for 3' to 5' DNA chain elongation. When. . .
CLM
       What is claimed is:
      . claim 24 wherein the water-solubilizing group is selected from the
       group consisting of sulfonate, phosphate, quaternary amine, sulfate,
       polyhydroxyl, and water-soluble polymer.
=> d his
     (FILE 'HOME' ENTERED AT 11:49:49 ON 04 JUL 2010)
     FILE 'BIOSIS, MEDLINE, CAPLUS, WPIDS, USPATFULL' ENTERED AT 11:50:31 ON
     04 JUL 2010
           4804 S (NUCLEOTIDE OR NUCLEOSIDE) AND BASE (4A) (LABEL? OR MARKER OR
L2
            531 S L1 AND LINKER (4A) (LABEL? OR MARKER OR DYE)
L3
             27 S L2 AND LINKER (6A) POLYMER
L4
              0 S L3 AND WATER SOLUBLE POLYMER
L5
             8 S L3 AND LINKER (4A) 50
1.6
              8 DUP REM L5 (0 DUPLICATES REMOVED)
             19 S L3 NOT L6
L7
L8
             19 DUP REM L7 (0 DUPLICATES REMOVED)
L9
             9 S L2 AND POLYMER (5A) WATER SOLUBLE
L10
             9 DUP REM L9 (0 DUPLICATES REMOVED)
=> d his
     (FILE 'HOME' ENTERED AT 11:49:49 ON 04 JUL 2010)
     FILE 'BIOSIS, MEDLINE, CAPLUS, WPIDS, USPATFULL' ENTERED AT 11:50:31 ON
     04 JUL 2010
L1
           4804 S (NUCLEOTIDE OR NUCLEOSIDE) AND BASE (4A) (LABEL? OR MARKER OR
L2
            531 S L1 AND LINKER (4A) (LABEL? OR MARKER OR DYE)
L3
             27 S L2 AND LINKER (6A) POLYMER
L4
             0 S L3 AND WATER SOLUBLE POLYMER
L5
             8 S L3 AND LINKER (4A) 50
L6
             8 DUP REM L5 (0 DUPLICATES REMOVED)
L7
             19 S L3 NOT L6
1.8
             19 DUP REM L7 (0 DUPLICATES REMOVED)
L9
             9 S L2 AND POLYMER (5A) WATER SOLUBLE
L10
              9 DUP REM L9 (0 DUPLICATES REMOVED)
=> s 12 and linker (4a)
MISSING TERM AFTER LINKER (4A
Operators must be followed by a search term, L-number, or query name.
=> s 12 and linker (4a) (peg or polyamide or polyphosphate or glycol or polyacetate
or poly?)
           117 L2 AND LINKER (4A) (PEG OR POLYAMIDE OR POLYPHOSPHATE OR GLYCOL
                OR POLYACETATE OR POLY?)
=> s 111 and soluble
           65 L11 AND SOLUBLE
```

```
=> s 112 not 13
L13
           44 L12 NOT L3
=> dup rem 113
PROCESSING COMPLETED FOR L13
            44 DUP REM L13 (0 DUPLICATES REMOVED)
L14
=> s 114 and 2003/pv
L15
             5 L14 AND 2003/PY
=> d 115 bib abs 1-5
L15 ANSWER 1 OF 5 USPATFULL on STN
AN
       2003:319498 USPATFULL
ΤТ
       Labeling reagents and labeled targets, target labeling processes and
       other processes for using same in nucleic acid determinations and
       analyses
TM
       Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES
       Rabbani, Elazar, New York, NY, UNITED STATES
ΡI
       US 20030225247
                          A1 20031204
                                                                    /--
       US 7166478
                          B2 20070123
                          A1 20020312 (10)
ΑI
      US 2002-96075
      Utility
      APPLICATION
FS
LREP
       ENZO LIFE SCIENCES, INC., c/o ENZO BIOCHEM, INC., 527 Madison Avenue,
       9th Floor, New York, NY, 10022
      Number of Claims: 286
CLMN
ECL
      Exemplary Claim: 1
DRWN
       15 Drawing Page(s)
LN.CNT 4499
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       This invention provides for labeling reagents, labeled targets and
       processes for preparing labeling reagents. The labeling reagents can
       take the form of cvanine dyes, xanthene dyes, porphyrin dyes, coumarin
       dyes or composite dyes. These labeling reagents are useful for labeling
      probes or targets, including nucleic acids and proteins. These reagents
       can be usefully applied to protein and nucleic acid probe based assays.
       They are also applicable to real-time detection processes.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L15 ANSWER 2 OF 5 USPATFULL on STN
AN
       2003:279097 USPATFULL
ΤI
       Releasable nonvolatile mass label molecules
IN
       Monforte, Joseph A., Berkeley, CA, United States
       Becker, Christopher H., Palo Alto, CA, United States
       Pollart, Daniel J., Menlo Park, CA, United States
       Shaler, Thomas A., Menlo Park, CA, United States
PA
       Sequenom Inc., San Diego, CA, United States (U.S. corporation)
PΙ
      US 6635452
                          B1 20031021
                                                                    <--
      US 1997-988024
ΑI
                               19971210 (8)
      US 1996-33037P
                               19961210 (60)
      US 1997-46719P
                               19970516 (60)
      Utility
FS
      GRANTED
EXNAM Primary Examiner: Riley, Jezia
LREP
      Heller Ehrman White & McAuliffe LLP, Seidman, Stephanie L.
CLMN
      Number of Claims: 90
ECI.
      Exemplary Claim: 1
DRWN 51 Drawing Figure(s); 35 Drawing Page(s)
LN.CNT 4660
```

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Using nonvolatile, releasable, mass-labels, the present invention provides for the synthesis and use of mass-labeled compounds to specifically interact with biomolecular targets. Following binding of the mass-labeled compounds to the target molecule, the unique mass-label can be analyzed using mass spectrometry to identify and characterize the target molecule. In one embodiment of the invention, a mass-labeled oligonucleotide probe is used to identify a specific gene sequence. A myriad of mass-labeled compounds may be produced for use in a wide variety of interactions such as oligonucleotide-oligonucleotide hybridization, polynucleotide-polynucleotide interactions, enzyme-substrate or substrate analog/intermediate interactions, polypeptide-nucleic acid interactions, protein-ligand interactions, receptor-ligand interactions, polypeptide-metal interactions, nucleic acid-metal interactions or antigen-antibody interactions. Also contemplated are combinatorial processes for creating large libraries of compounds permitting rapid screening for a wide variety of targets.

#### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

```
L15 ANSWER 3 OF 5 USPATFULL on STN
```

AN 2003:273357 USPATFULL

TI Manipulation of microparticles in microfluidic systems

IN Mehta, Tammy Burd, San Jose, CA, United States Kopf-Sill, Anne R., Portola Valley, CA, United States

Parce, J. Wallace, Palo Alto, CA, United States Chow, Andrea W., Los Altos, CA, United States Bousse, Luc J., Los Altos, CA, United States

Knapp, Michael R., Redwood City, CA, United States Nikiforov, Theo T., San Jose, CA, United States

Gallagher, Steve, Palo Alto, CA, United States
PA Caliper Technologies Corp., Mountain View, CA, United States (U.S.

corporation)
PI US 6632655 B1 20031014 <--

PI US 6632655 B1 20031014 AI US 2000-510626 20000222 (9) PRAI US 1999-128643P 19990409 (60) US 1999-127825P 19990405 (60) US 1999-121223P 19990223 (60)

DT Utility FS GRANTED

EXNAM Primary Examiner: Ponnaluri, Padmashri; Assistant Examiner: Tran, My

LREP Quine Intellectual Property Law Group, P.C., Murphy, Matthew B., McKenna, Donald R.

CLMN Number of Claims: 71 ECL Exemplary Claim: 1

DRWN 28 Drawing Figure(s); 19 Drawing Page(s)

LN.CNT 4515

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Arrays of flowable or fixed particle sets are used in microfluidic systems for performing assays and modifying hydrodynamic flow. Also provided are assays utilizing flowable or fixed particle sets within a microfluidic system, as well as kits, apparatus and integrated systems comprising arrays and array members.

## CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L15 ANSWER 4 OF 5 USPATFULL on STN

AN 2003:237696 USPATFULL

TI Antisense imaging of gene expression of the brain in vivo

IN Partridge, William M., Pacific Palisades, CA, UNITED STATES

```
Boado, Ruben J., Agoura Hills, CA, UNITED STATES
       The Regents of the University of California Office of Technology
PA
       Transfer (U.S. corporation)
       US 20030165853
                          A1 20030904
PT
                                                                    <--
       US 2001-5996
                          A1 20011203 (10)
AΙ
PRAI
       US 2000-250990P
                               20001204 (60)
       Utility
       APPLICATION
       OUINE INTELLECTUAL PROPERTY LAW GROUP, P.C., P O BOX 458, ALAMEDA, CA,
LREP
       94501
CLMN
     Number of Claims: 61
ECL
       Exemplary Claim: 1
DRWN 12 Drawing Page(s)
LN.CNT 2661
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AR
       This invention provides imaging reagents for the detection of a gene or
       gene expression product (e.g. mRNA) in a brain cell in vivo. Preferred
       reagents comprise a detectable label attached to a first nucleic acid
       that specifically hybridizes to the gene or to a nucleic acid
       transcribed from the gene. The first nucleic acid is linked to a
       targeting ligand that is capable of binding a receptor on a cell
       comprising the blood brain barrier and crossing said blood brain
       barrier.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L15 ANSWER 5 OF 5 USPATFULL on STN
       2003:30278 USPATFULL
AN
ΤI
       Releasable nonvolatile mass-label molecules
TN
       Monforte, Joseph A., Berkeley, CA, UNITED STATES
       Becker, Christopher H., Palo Alto, CA, UNITED STATES
       Pollart, Daniel J., Menlo Park, CA, UNITED STATES
       Shaler, Thomas A., Menlo Park, CA, UNITED STATES
       US 20030022225
                          A1 20030130
PΙ
                                                                    <--
       US 7132519
                          B2 20061107
AΤ
       US 2002-202189
                          A1 20020722 (10)
RLI
       Continuation of Ser. No. US 1997-988024, filed on 10 Dec 1997, PENDING
PRAI
       US 1996-33037P
                               19961210 (60)
       US 1997-46719P
                               19970516 (60)
DT
      Utility
FS
       APPLICATION
      Stephanie Seidman, Heller Ehrman White & McAuliffe LLP, 7th Floor, 4350
       La Jolla Village Drive, San Diego, CA, 92122
CLMN
       Number of Claims: 122
ECL
       Exemplary Claim: 1
DRWN
      35 Drawing Page(s)
IN.CNT 4085
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AB
       Releasable tag reagents for use in the detection and analysis of target
       molecules, particular in mass spectrometric analyses are provided. Also
       provided are methods of detection that employ releasable tag reagents.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
=> s 114 not 115
```

L16

39 L14 NOT L15

L16 ANSWER 1 OF 39 USPATFULL on STN

=> d 116 bib abs 1-39

```
AN
       2010:77816 USPATFULL
TT
       POLYNUCLEOTIDES AND RELATED NANOASSEMBLIES, STRUCTURES, ARRANGEMENTS,
       METHODS AND SYSTEMS
      MAUNE, Hareem T., Pasadena, CA, UNITED STATES
       Han, Si-Ping, Yorba Linda, CA, UNITED STATES
       Barish, Robert D., Pasadena, CA, UNITED STATES
       Bockrath, Marc W., Diamond Bar, CA, UNITED STATES
       Goddard, William A., Pasadena, CA, UNITED STATES
       Rothemund, Paul W.K., Pasadena, CA, UNITED STATES
       Winfree, Erik, Altadena, CA, UNITED STATES
PΙ
      US 20100069621
                          A1 20100318
                          A1 20090812 (12)
AΙ
      US 2009-540052
PRAI
      US 2008-188854P
                               20080813 (61)
      US 2008-189792P
                               20080822 (61)
      US 2009-170564P
                              20090417 (61)
DТ
      Utility
FS
      APPLICATION
LREP
      Steinfl & Bruno, 301 N Lake Ave Ste 810, Pasadena, CA, 91101, US
CLMN Number of Claims: 20
ECL
      Exemplary Claim: 1
       22 Drawing Page(s)
DRWN
LN.CNT 2929
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       A linker polynucleotide for attaching a nanomaterial
       to a polynucleotidic platform and related nanoassemblies, arrangements,
       structures, methods and systems.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L16 ANSWER 2 OF 39 USPATFULL on STN
       2009:363316 USPATFULL
AN
       CIS REACTIVE OXYGEN QUENCHERS INTEGRATED INTO LINKERS
TI
       Otto, Geoffrey, San Carlos, CA, UNITED STATES
       Shen, Gene, Santa Clara, CA, UNITED STATES
       Kong, Xiangxu, Foster City, CA, UNITED STATES
       Emig, Robin, Belmont, CA, UNITED STATES
PA
      Pacific Biosciences of California, Inc., Menlo Park, CA, UNITED STATES
       (U.S. corporation)
PΙ
      US 20090325260
                           A1 20091231
ΑI
      US 2009-367411
                          A1 20090206 (12)
                               20080207 (61)
PRAI
      US 2008-26992P
DT
      Utility
FS
      APPLICATION
LREP
      MORGAN, LEWIS & BOCKIUS LLP (SF), One Market, Spear Street Tower, Suite
       2800, San Francisco, CA, 94105, US
CLMN
      Number of Claims: 35
ECI.
      Exemplary Claim: 1
      12 Drawing Page(s)
DRWN
LN.CNT 1318
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AB
       The present invention provides methods and compositions for performing
       illuminated reactions, particularly sequencing reactions, while
       mitigating and/or preventing photodamage to reactants that can result
       from prolonged illumination. In particular, the invention provides
       methods and compositions for incorporating photoprotective agents into
       conjugates comprising reporter molecules and nucleoside
       polyphosphates.
```

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L16 ANSWER 3 OF 39 USPATFULL on STN

```
AN
       2009:348386 USPATFULL
TT
       BIOASSAY SYSTEM INCLUDING OPTICAL DETECTION APPARATUSES, AND METHOD FOR
       DETECTING BIOMOLECULES
TN
       CHIOU, CHUNG-FAN, Cyonglin Township, TAIWAN, PROVINCE OF CHINA
       Chu, Cheng-Wei, Yonghe City, TAIWAN, PROVINCE OF CHINA
       Li, Yu-Tang, Tucheng City, TAIWAN, PROVINCE OF CHINA
       Chu, Chang-Sheng, Hsinchu City, TAIWAN, PROVINCE OF CHINA
       Chung, Shuang-Chao, Jhongli City, TAIWAN, PROVINCE OF CHINA
       Fan, Chih-Hsun, Hsinchu Citv, TAIWAN, PROVINCE OF CHINA
       Industrial Technology Research Institute (non-U.S. corporation)
PΙ
       US 20090311774
                          A1 20091217
AΙ
      US 2009-500567
                          A1 20090709 (12)
RLI
      Continuation-in-part of Ser. No. US 2008-255044, filed on 21 Oct 2008,
       PENDING
PRAT
      US 2007-996016P
                               20071025 (60)
      US 2008-36652P
                              20080314 (61)
DТ
      Utility
FS
      APPLICATION
LREP
       FINNEGAN, HENDERSON, FARABOW, GARRETT & DUNNER, LLP, 901 NEW YORK
      AVENUE, NW, WASHINGTON, DC, 20001-4413, US
CLMN
      Number of Claims: 25
ECL
       Exemplary Claim: 1
DRWN
       7 Drawing Page(s)
LN.CNT 1674
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       A bioassay system is disclosed. The bioassay system may include a
       plurality of optical detection apparatuses, each of which includes a
       substrate having a light detector, and a linker site formed over the
       light detector, the linker site being treated to affix the biomolecule
       to the linker site. The linker site is proximate to the light detector
       and is spaced apart from the light detector by a distance of less than
       or equal to 100 micrometers. The light detector collects light emitted
       from the biomolecule within a solid angle of greater than or equal to
       0.8 SI steridian. The optical detection apparatus may further include an
       excitation light source formed over the substrate so as to provide a
       light source for exciting a fluorophore attached to the biomolecule.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L16 ANSWER 4 OF 39 USPATFULL on STN
AN
       2009:213280 USPATFULL
ΤI
       Chase Ligation Sequencing
IN
       Hendrickson, Cynthia, Wenham, MA, UNITED STATES
PA
      Applied Biosystems Inc., Foster City, CA, UNITED STATES (U.S.
       corporation)
PΤ
      US 20090191553
                          A1 20090730
      US 2008-243925
                          A1 20081001 (12)
ΑI
      US 2007-976757P
                               20071001 (60)
PRAI
DT
      Utility
FS
      APPLICATION
LREP
      MILA KASAN, PATENT DEPT., APPLIED BIOSYSTEMS, 850 LINCOLN CENTRE DRIVE,
       FOSTER CITY, CA, 94404, US
CLMN
      Number of Claims: 44
ECL
      Exemplary Claim: 1
      70 Drawing Page(s)
LN.CNT 7177
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
```

AB In various embodiments, the present teachings provide sequencing methods which facilitate enhancing the efficiency of ligation and/or increasing sequencing reads. Various embodiments of the methods enable sequencing through template regions for which complementary labeled extension

probes are unavailable or insufficient. In various embodiments, one or more rounds of ligation with unlabeled extension probes can be used maddition to a round of ligation with labeled extension probe. In various embodiments, for example, such methods can facilitate extension on template polynucleotides that do not bind labeled extension probe in the first round of ligation.

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CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L16 ANSWER 5 OF 39 USPATFULL on STN
AN
       2009:162778 USPATFULL
ΤI
       BIOASSAY SYSTEM INCLUDING OPTICAL DETECTION APPARATUSES, AND METHOD FOR
       DETECTING BIOMOLECULES
TN
       CHIOU, Chung-Fan, Cyonglin Township, TAIWAN, PROVINCE OF CHINA
       CHU, Cheng-Wei, Yonghe City, TAIWAN, PROVINCE OF CHINA
       CHANG, Shang-Chia, Zhubei City, TAIWAN, PROVINCE OF CHINA
       LI, Yu-Tang, Tucheng City, TAIWAN, PROVINCE OF CHINA
       PAN, Chao-Chi, Hsinchu City, TAIWAN, PROVINCE OF CHINA
       YAO, Bin-Cheng, Taipei City, TAIWAN, PROVINCE OF CHINA
PΙ
       US 20090146076
                          A1 20090611
      US 2008-255044
                           A1 20081021 (12)
ΑI
PRAI
      US 2007-996016P
                               20071025 (60)
      US 2008-36652P
                               20080314 (61)
DT
      Utility
FS
       APPLICATION
LREP
       FINNEGAN, HENDERSON, FARABOW, GARRETT & DUNNER, LLP, 901 NEW YORK
      AVENUE, NW, WASHINGTON, DC, 20001-4413, US
CLMN
      Number of Claims: 61
ECL
      Exemplary Claim: 1
DRWN
      6 Drawing Page(s)
LN.CNT 1705
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       A bioassay system is disclosed. The bioassay system may include a
```

A bioassay system is disclosed. The bioassay system may include a plurality of optical detection apparatuses, each of which includes a substrate having a light detector, and a linker site formed over the light detector, the linker site being treated to affix the biomolecule to the linker site. The linker site is proximate to the light detector and is spaced apart from the light detector by a distance of less than or equal to 100 micrometers. The light detector collects light emitted from the biomolecule within a solid angle of greater than or equal to 0.8 SI steridian. The optical detection apparatus may further include an excitation light source formed over the substrate so as to provide a

light source for exciting a fluorophore attached to the biomolecule.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

```
L16 ANSWER 6 OF 39 USPATFULL on STN
       2009:152569 USPATFULL
AN
       Manipulation of Microparticles In Microfluidic Systems
ΤI
IN
       Mehta, Tammy Burd, San Jose, CA, UNITED STATES
       Kopf-Sill, Anne R., Portola Valley, CA, UNITED STATES
       Parce, J. Wallace, Palo Alto, CA, UNITED STATES
       Chow, Andrea W., Los Altos, CA, UNITED STATES
       Bousse, Luc J., Los Altos, CA, UNITED STATES
       Knapp, Michael R., Palo Alto, CA, UNITED STATES
       Nikiforov, Theo T., San Jose, CA, UNITED STATES
       Gallagher, Steve, Palo Alto, CA, UNITED STATES
PΆ
      CALIPER LIFE SCIENCES, INC., Mountain View, CA, UNITED STATES (U.S.
       corporation)
PT
      US 20090137413
                        A1 20090528
                         A1 20071030 (11)
AΤ
      US 2007-928808
```

```
RLT
      Division of Ser. No. US 2003-606201, filed on 25 Jun 2003, PENDING
       Continuation of Ser. No. US 2000-510626, filed on 22 Feb 2000, Pat. No.
      US 6632655
      WO 2000-US4486
                              20000222
PRAT
       WO 2000-US4522
                              20000222
      US 1999-121223P
                              19990223 (60)
      US 1999-127825P
                              19990405 (60)
      US 1999-128643P
                              19990409 (60)
      Utility
      APPLICATION
FS
      CARDINAL LAW GROUP, Caliper Life Sciences, Inc., 1603 Orrington Avenue,
      Suite 2000, Evanston, IL, 60201, US
CLMN
     Number of Claims: 5
ECL
      Exemplary Claim: 1
DRWN
     19 Drawing Page(s)
LN.CNT 4104
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
      Arrays of flowable or fixed particle sets are used in microfluidic
       systems for performing assays and modifying hydrodynamic flow. Also
       provided are assays utilizing flowable or fixed particle sets within a
       microfluidic system, as well as kits, apparatus and integrated systems
      comprising arrays and array members.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L16 ANSWER 7 OF 39 USPATFULL on STN
       2009:145929 USPATFULL
       SIALIC ACID ABC TRANSPORTERS IN PROKARYOTES THERAPEUTIC TARGETS
IN
       Gibson, Bradford W., Berkeley, CA, UNITED STATES
       Munson, Robert S., Hilliard, OH, UNITED STATES
       Post, Deborah M., Fairfax, CA, UNITED STATES
PA
       BUCK INSTITUTE, Novato, CA, UNITED STATES (U.S. corporation)
ΡI
      US 20090131524
                         A1 20090521
      US 2006-916975
                          A1 20060531 (11)
ΑI
      WO 2006-US21202
                              20060531
                              20081222 PCT 371 date
PRAI
      US 2005-689151P
                              20050607 (60)
DT
      Utility
FS
      APPLICATION
LREP
      Weaver Austin Villeneuve & Sampson LLP, P.O. BOX 70250, OAKLAND, CA,
      94612-0250, US
CLMN
      Number of Claims: 39
ECL
      Exemplary Claim: 1
DRWN
     5 Drawing Page(s)
LN.CNT 2903
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       This invention provides a novel bacterial sialic acid transporter that
AR
       is a member of the family of ABC transporters. The transporter is a
       useful target for pharmaceuticals.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L16 ANSWER 8 OF 39 USPATFULL on STN
AN
       2009:76424 USPATFULL
ΤI
       Label target and labeling reagents comprising rigid group backbones
TM
       Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES
```

Rabbani, Elazar, New York, NY, UNITED STATES

A1 20040122 (10)

STATES (U.S. corporation)

US 2004-763102

US 20090069500 A1 20090312

Enzo Life Sciences, Inc., c/o Enzo Biochem, Inc., New York, NY, UNITED

PΑ

PT

AΤ

```
RLT
       Division of Ser. No. US 2002-96075, filed on 12 Mar 2002, Pat. No. US
       7166478
       Utility
       APPLICATION
LREP
       ENZO BIOCHEM, INC., 527 MADISON AVENUE (9TH FLOOR), NEW YORK, NY, 10022,
CLMN
       Number of Claims: 77
ECL
       Exemplary Claim: 1-286
DRWN
       15 Drawing Page(s)
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       This invention provides for labeling reagents, labeled targets and
       processes for preparing labeling reagents. The labeling reagents can
       take the form of cyanine dyes, xanthene dyes, porphyrin dyes, coumarin
       dyes or composite dyes. These labeling reagents are useful for labeling
       probes or targets, including nucleic acids and proteins. These reagents
       can be usefully applied to protein and nucleic acid probe based assays.
       They are also applicable to real-time detection processes.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L16 ANSWER 9 OF 39 USPATFULL on STN
       2008:305957 USPATFULL
AN
ΤI
       DETECTABLE LABELED NUCLEOSIDE ANALOGS AND METHODS OF USE
       THEREOF
       Bodepudi, Veeraiah, San Ramon, CA, UNITED STATES
       GUPTA, Amar, Danville, CA, UNITED STATES
       Will, Stephen G., Oakland, CA, UNITED STATES
PT
       US 20080268441
                          A1 20081030
       US 7452674
                          B2 20081118
AΙ
       US 2007-742097
                          A1 20070430 (11)
DT
       Utility
FS
       APPLICATION
LREP
       Roche Molecular Systems, Inc., Patent Law Department, 4300 Hacienda
       Drive, Pleasanton, CA, 94588, US
CLMN
     Number of Claims: 8
ECL
       Exemplary Claim: 1-68
DRWN
      No Drawings
LN.CNT 2148
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       The invention relates to detectable labels useful for detection of
       nucleotide sequences. Specifically, the invention relates to
       labeled-imidazole-PEG compounds, such as nucleosides, nucleotides, and
       nucleic acids incorporating such compounds, and methods utilizing such
       compounds. The invention further relates to kits comprising labeled
       imidazole-PEG compounds.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L16 ANSWER 10 OF 39 USPATFULL on STN
       2008:238073 USPATFULL
ΑN
       POLYNUCLEIC ACID-ATTACHED PARTICLES AND THEIR USE IN GENOMIC ANALYSIS
ΤI
IN
       Loge, Gary W., State College, PA, UNITED STATES
PA
       LCM Technologies, Inc., State College, PA, UNITED STATES (U.S.
       corporation)
PΙ
       US 20080206758
                          A1 20080828
       US 2007-872892
                          A1 20071016 (11)
AΤ
PRAT
      US 2006-829719P
                               20061017 (60)
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WOODCOCK WASHBURN LLP, CIRA CENTRE, 12TH FLOOR, 2929 ARCH STREET,

DT

FS

Utility

APPLICATION

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PHILADELPHIA, PA, 19104-2891, US
      Number of Claims: 25
CLMN
      Exemplary Claim: 1
ECL
DRWN
      8 Drawing Page(s)
LN.CNT 1190
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       Disclosed are methods for preparing particle-linked polynucleotides, and
       using the particle linked polynucleotides in genomic analysis. The
       particles as disclosed are characterized as having a size variance of
       less than 2%.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L16 ANSWER 11 OF 39 USPATFULL on STN
AN
       2007:237793 USPATFULL
ТΤ
       DETECTABLE LABELED NUCLEOSIDE ANALOGS AND METHODS OF USE
       THEREOF
TM
       Bodepudi, Veeraiah, San Ramon, CA, UNITED STATES
       Gupta, Amar, Danville, CA, UNITED STATES
       Will, Stephen G., Oakland, CA, UNITED STATES
       ROCHE MOLECULAR SYSTEMS, INC., Alameda, CA, UNITED STATES (U.S.
PA
       corporation)
       US 20070208169
PΙ
                          A1 20070906
      US 7456266
                          B2 20081125
       US 2007-742070
                          A1
                              20070430 (11)
AΙ
       Division of Ser. No. US 2003-719257, filed on 21 Nov 2003, GRANTED, Pat.
RI.T
PRAI
      US 2002-428484P
                             20021122 (60)
DT
      Utility
FS
      APPLICATION
LREP
      ROCHE MOLECULAR SYSTEMS INC, PATENT LAW DEPARTMENT, 1145 ATLANTIC
      AVENUE, ALAMEDA, CA, 94501, US
CLMN
      Number of Claims: 22
ECL
      Exemplary Claim: 1
DRWN
      No Drawings
LN.CNT 2068
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
      The invention relates to detectable labels useful for detection of
       nucleotide sequences. Specifically, the invention relates to
       labeled-imidazole-PEG compounds, such as nucleosides, nucleotides, and
       nucleic acids incorporating such compounds, and methods utilizing such
       compounds. The invention further relates to kits comprising labeled
       imidazole-PEG compounds.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L16 ANSWER 12 OF 39 USPATFULL on STN
       2007:231302 USPATFULL
AN
       DETECTABLE LABELED NUCLEOSIDE ANALOGS AND METHODS OF USE
ΤI
       THEREOF
       Bodepudi, Veeraiah, San Ramon, CA, UNITED STATES
       Gupta, Amar, Danville, CA, UNITED STATES
       Will, Stephen G., Oakland, CA, UNITED STATES
PA
      ROCHE MOLECULAR SYSTEMS, INC., Alameda, CA, UNITED STATES (U.S.
       corporation)
      US 20070202576
                          A1 20070830
       US 7501504
                          B2 20090310
      US 2007-742123
                         A1 20070430 (11)
AΤ
RLT
      Division of Ser. No. US 2003-719257, filed on 21 Nov 2003, GRANTED, Pat.
      No. US 7220847
PRAI US 2002-428484P
                           20021122 (60)
```

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DT
     Utility
FS
       APPLICATION
       ROCHE MOLECULAR SYSTEMS INC, PATENT LAW DEPARTMENT, 1145 ATLANTIC
LREP
       AVENUE, ALAMEDA, CA, 94501, US
CLMN Number of Claims: 16
ECL Exemplary Claim: 1-77
DRWN No Drawings
LN.CNT 2056
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       The invention relates to detectable labels useful for detection of
       nucleotide sequences. Specifically, the invention relates to
       labeled-imidazole-PEG compounds, such as nucleosides, nucleotides, and
       nucleic acids incorporating such compounds, and methods utilizing such
       compounds. The invention further relates to kits comprising labeled
       imidazole-PEG compounds.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L16 ANSWER 13 OF 39 USPATFULL on STN
AN
       2007:68444 USPATFULL
ΤI
       "Met/fret based method of target nucleic acid detection whereby the
       donor/acceptor moieties are on complementary strands"
       Islam, Amirul, Hyderabad, INDIA
       Hazea, Papia, Hyderabed, INDIA
       US 20070059690
                        A1 20070315
       US 2003-516361
                          A1 20030530 (10)
AΤ
       WO 2003-TN204
                               20030530
                               20041130 PCT 371 date
PRAI
       IN 2002-MU487
                               20020531
       Utility
DT
FS
       APPLICATION
LREP
       STEPHAN A. PENDORF, P.A., PENDORF & CUTLIFF, 5111 MEMORIAL HIGHWAY,
       TAMPA, FL, 33634, US
CLMN
     Number of Claims: 57
ECL
      Exemplary Claim: 1-62
DRWN
      38 Drawing Page(s)
LN.CNT 3233
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       Disclosure of a method for the detection and quantitation of
       polynucleotide sequences in a sample of biological or non-biological
       material through target poly nucleotide sequence amplification
       by polymerase chain reaction using chemically labeled oligonucleotide
       amplification primers and formation of an entity between the amplified
       polynucleotide sequence and chemically labeled polynucleotide having a
       sequence complementary to the target polynucleotide sequence for
       determining the identity and/or presence and/or quantitation of the
       target poly nucleotide sequences. The chemical label
       covalently attached to the oligonucleotide amplification primer and
       polynucleotide or oligonucleotide comprise molecular energy transfer
       labels (donor and acceptor). It is again a very sensitive, rapid and
       reliable method with better sensitivity, specificity and reliability for
       the detection of polynucleotide sequence. It also greatly reduces the
```

L16 ANSWER 14 OF 39 USPATFULL on STN

AN 2006:202424 USPATFULL

measurements.

TI Labeling reagents and labeled targets comprising nonmetallic porphyrins

possibility of amplification product carry-over contamination and adaptable for many formats of nucleic acids amplifications and real time

```
ΤN
       Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES
```

Rabbani, Elazar, New York, NY, UNITED STATES

PA Enzo Life Sciences, Inc., c/o Enzo Biochem, Inc., New York, NY, UNITED STATES (U.S. corporation)

PT US 20060172308 A1 20060803 US 7537751 B2 20090526

US 2004-763088 AΙ A1 20040122 (10) Division of Ser. No. US 2002-96075, filed on 12 Mar 2002, PENDING

RLI

Utility DT

FS APPLICATION

LREP ENZO BIOCHEM, INC., 527 MADISON AVENUE (9TH FLOOR), NEW YORK, NY, 10022,

CLMN Number of Claims: 19

ECL Exemplary Claim: 1

DRWN 15 Drawing Page(s) LN.CNT 3541

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

This invention provides for labeling reagents, labeled targets and processes for preparing labeling reagents. The labeling reagents can take the form of cyanine dyes, xanthene dyes, porphyrin dyes, coumarin dyes or composite dyes. These labeling reagents are useful for labeling probes or targets, including nucleic acids and proteins. These reagents can be usefully applied to protein and nucleic acid probe based assays. They are also applicable to real-time detection processes.

# CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L16 ANSWER 15 OF 39 USPATFULL on STN

AN 2006:152267 USPATFULL

Antagonizing an adenosine A2A receptor to ameliorate one or more TI components of addictive behavior

Diamond, Ivan F., Berkeley, CA, UNITED STATES

Gordon, Adrienne S., Kensington, CA, UNITED STATES

The Regents of the University of California (U.S. corporation) PA A1 20060615 PI US 20060128708

ΑI US 2005-153725 A1 20050614 (11)

PRAI US 2004-581143P 20040617 (60)

DT Utility

FS APPLICATION

LREP QUINE INTELLECTUAL PROPERTY LAW GROUP, P.C., P O BOX 458, ALAMEDA, CA, 94501, US

CLMN Number of Claims: 50

ECL Exemplary Claim: 1

DRWN 6 Drawing Page(s)

LN.CNT 2235

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

This invention provides a method of mitigating/ameliorating one or more AR components of addictive behavior associated with chronic consumption of a substance of abuse, or withdrawal therefrom. The method typically involves administering to a subject in need thereof an adenosine A2A receptor antagonist in an amount sufficient to ameliorate said one or more components of addictive behavior.

# CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L16 ANSWER 16 OF 39 USPATFULL on STN

AN 2006:46815 USPATFULL

Bio-barcode based detection of target analytes

TN Mirkin, Chad A., Wilmette, IL, UNITED STATES Nam, Jwa-Min, Berkeley, CA, UNITED STATES Oh, Byung-Keun, Evanston, IL, UNITED STATES

```
Thaxton, C. Shad, Chicago, IL, UNITED STATES
       Georganopoulou, Dimitra, Chicago, IL, UNITED STATES
PA
       Nanosphere, Inc. (U.S. corporation)
PT
       US 20060040286
                           A1 20060223
AΙ
       US 2005-127808
                           A1 20050512 (11)
RLI
       Continuation-in-part of Ser. No. US 2004-877750, filed on 25 Jun 2004,
       PENDING Continuation-in-part of Ser. No. WO 2004-US20493, filed on 25
       Jun 2004, PENDING Continuation-in-part of Ser. No. US 2002-108211, filed
       on 27 Mar 2002, GRANTED, Pat. No. US 6974669 Continuation-in-part of
       Ser. No. US 2001-820279, filed on 28 Mar 2001, GRANTED, Pat. No. US
       6750016
PRAI
       US 2004-570723P
                               20040512 (60)
       US 2004-585294P
                               20040701 (60)
       US 2005-645455P
                               20050119 (60)
       US 2003-506708P
                               20030926 (60)
       US 2003-482979P
                               20030627 (60)
       US 2003-496893P
                               20030821 (60)
                               20031028 (60)
       US 2003-515243P
                               20031218 (60)
       US 2003-530797P
       US 2001-350560P
                               20011113 (60)
       Utility
FS
       APPLICATION
       MCDONNELL BOEHNEN HULBERT & BERGHOFF LLP, 300 S. WACKER DRIVE, 32ND
LREP
       FLOOR, CHICAGO, IL, 60606, US
CLMN
       Number of Claims: 50
       Exemplary Claim: 1
ECL
DRWN
       27 Drawing Page(s)
LN.CNT 4753
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       The present invention relates to screening methods, compositions, and
AB
       kits for detecting for the presence or absence of one or more target
       analytes, e.g. biomolecules, in a sample. In particular, the present
       invention relates to a method that utilizes reporter oligonucleotides as
       biochemical barcodes for detecting multiple protein structures or other
       target analytes in a solution.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L16 ANSWER 17 OF 39 USPATFULL on STN
ΑN
       2006:40616 USPATFULL
       Processes for incorporating nucleic acid sequences into an analyte or
TI
       library of analytes
IN
       Rabbani, Elazar, New York, NY, UNITED STATES
       Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES
       Donegan, James J., Long Beach, NY, UNITED STATES
       Coleman, Jack, East Northport, NY, UNITED STATES
       Liu, Dakai, Islip, NY, UNITED STATES
PA
       Enzo Life Sciences, Inc., New York, NY, UNITED STATES (U.S. corporation)
                          A1 20060216
ΡI
       US 20060035264
       US 2005-237466
                           A1 20050927 (11)
AΙ
RLI
       Division of Ser. No. US 2002-96076, filed on 12 Mar 2002, PENDING
       Utility
FS
       APPLICATION
LREP
       ENZO BIOCHEM, INC., 527 MADISON AVENUE (9TH FLOOR), NEW YORK, NY, 10022,
CLMN
       Number of Claims: 69
ECL
       Exemplary Claim: 1-413
DRWN
       15 Drawing Page(s)
LN.CNT 4099
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AR
       This invention provides for compositions for use in real time nucleic
```

acid detection processes. Such real time nucleic acid detection processes are carried out with energy transfer elements attached to nucleic acid primers, nucleotides, nucleic acid probes or nucleic acid binding agents. Real time nucleic acid detection allows for the qualitative or quantitative detection or determination of single-stranded or double-stranded nucleic acids of interest in a sample. Other processes are provided by this invention including processes for removing a portion of a homopolymeric sequence, e.g., poly A sequence or tail, from an analyte or library of analytes. Compositions useful in carrying out such removal processes are also described and provided.

# CAS INDEXING IS AVAILABLE FOR THIS PATENT.

- L16 ANSWER 18 OF 39 USPATFULL on STN
- AN 2006:34199 USPATFULL
- TI Processes for quantitative or qualitative detection of single-stranded or double-stranded nucleic acids
- IN Rabbani, Elazar, New York, NY, UNITED STATES Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES Donegan, James J., Long Beach, NY, UNITED STATES Coleman, Jack, East Northport, NY, UNITED STATES

A1 20060209

- PI US 20060029968
- AI US 2005-235516 A1 20050926 (11)

Liu, Dakai, Islip, NY, UNITED STATES

- RLI Division of Ser. No. US 2002-96076, filed on 12 Mar 2002, PENDING
- DT Utility
- FS APPLICATION
- LREP ENZO BIOCHEM, INC., 527 MADISON AVENUE (9TH FLOOR), NEW YORK, NY, 10022,
- CLMN Number of Claims: 275
- ECL Exemplary Claim: 1-33
- DRWN 15 Drawing Page(s)
- LN.CNT 5182
- CAS INDEXING IS AVAILABLE FOR THIS PATENT.
- AB This invention provides for compositions for use in real time nucleic acid detection processes. Such real time nucleic acid detection processes are carried out with energy transfer elements attached to nucleic acid primers, nucleotides, nucleic acid probes or nucleic acid binding agents. Real time nucleic acid detection allows for the qualitative or quantitative detection or determination of single-stranded or double-stranded nucleic acids of interest in a sample. Other processes are provided by this invention including processes for removing a portion of a homopolymeric sequence, e.g., poly A sequence or tail, from an analyte or library of analytes. Compositions useful in carrying out such removal processes are also described and provided.

## CAS INDEXING IS AVAILABLE FOR THIS PATENT.

- L16 ANSWER 19 OF 39 USPATFULL on STN
- AN 2006:27907 USPATFULL
- TI Site- or sequence-specific process for cleaving analytes and library of analytes
- IN Rabbani, Elazar, New York, NY, UNITED STATES
  Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES
  Donegan, James J., Long Beach, NY, UNITED STATES
  Coleman, Jack, East Northport, NY, UNITED STATES
  Liu, Dakai, Islip, NY, UNITED STATES
- PA Enzo Life Sciences, Inc., New York, NY, UNITED STATES (U.S. corporation)
- PI US 20060024738 A1 20060202

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US 2005-237467
                          A1 20050927 (11)
AΤ
RLT
       Division of Ser. No. US 2002-96076, filed on 12 Mar 2002, PENDING
DT
      Utility
FS
      APPLICATION
LREP
       ENZO BIOCHEM, INC., 527 MADISON AVENUE (9TH FLOOR), NEW YORK, NY, 10022,
CLMN
      Number of Claims: 555
ECL
      Exemplary Claim: 1
DRWN
      15 Drawing Page(s)
LN.CNT 6144
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       This invention provides for compositions for use in real time nucleic
       acid detection processes. Such real time nucleic acid detection
       processes are carried out with energy transfer elements attached to
       nucleic acid primers, nucleotides, nucleic acid probes or nucleic acid
       binding agents. Real time nucleic acid detection allows for the
       qualitative or quantitative detection or determination of
       single-stranded or double-stranded nucleic acids of interest in a
       sample. Other processes are provided by this invention including
       processes for removing a portion of a homopolymeric sequence, e.g., poly
       A sequence or tail, from an analyte or library of analytes. Compositions
       useful in carrying out such removal processes are also described and
       provided.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L16 ANSWER 20 OF 39 USPATFULL on STN
AN
       2006:27906 USPATFULL
       Process for removal of homopolymeric sequence portion from analyte(s)
TI
       and library of analytes
       Babbani, Elazar, New york, NY, UNITED STATES
       Stavrianopoulos, Jannis G., Baysnore, NY, UNITED STATES
       Donegan, James J., Long Beach, NY, UNITED STATES
       Coleman, Jack, East Northport, NY, UNITED STATES
       Liu, Dakai, Islip, NY, UNITED STATES
PA
       Enzo Life Sciences, Inc., New York, NY, UNITED STATES (U.S. corporation)
PΙ
      US 20060024737
                           A1 20060202
      US 7550265
                           B2 20090623
ΑI
      US 2005-237442
                          A1 20050927 (11)
      Division of Ser. No. US 2002-96076, filed on 12 Mar 2002, PENDING
RI.T
DT
      Utility
FS
      APPLICATION
LREP
      ENZO BIOCHEM, INC., 527 MADISON AVENUE (9TH FLOOR), NEW YORK, NY, 10022,
CLMN
      Number of Claims: 17
ECI.
      Exemplary Claim: 1-527
DRWN
      15 Drawing Page(s)
LN.CNT 3943
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AB
       This invention provides for compositions for use in real time nucleic
       acid detection processes. Such real time nucleic acid detection
       processes are carried out with energy transfer elements attached to
       nucleic acid primers, nucleotides, nucleic acid probes or nucleic acid
       binding agents. Real time nucleic acid detection allows for the
       qualitative or quantitative detection or determination of
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single-stranded or double-stranded nucleic acids of interest in a sample. Other processes are provided by this invention including processes for removing a portion of a homopolymeric sequence, e.g., poly A sequence or tail, from an analyte or library of analytes. Compositions useful in carrying out such removal processes are also described and

B2 20080708

US 7396647

provided.

# CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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L16 ANSWER 21 OF 39 USPATFULL on STN
```

AN 2006:27904 USPATFULL

TΙ Chimeric nucleic acid constructs and compositions comprising sets of nucleic acid constructs

IN Rabbani, Elazar, New York, NY, UNITED STATES Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES Donegan, James J., Long Beach, NY, UNITED STATES Coleman, Jack, East Northport, NY, UNITED STATES

Liu, Dakai, Lslip, NY, UNITED STATES PA Enzo Life Sciences, Inc., New York, NY, UNITED STATES (U.S. corporation)

PΙ US 20060024735 A1 20060202 US 7547772 B2 20090616

US 2005-236151 A1 20050927 (11)

RLI Division of Ser. No. US 2002-96076, filed on 12 Mar 2002, PENDING

DT Utility

ΑI

FS APPLICATION

LREP ENZO BIOCHEM, INC., 527 MADISON AVENUE (9TH FLOOR), NEW YORK, NY, 10022,

CLMN Number of Claims: 52

ECL Exemplary Claim: 1-404 DRWN 15 Drawing Page(s)

LN.CNT 4013

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

This invention provides for compositions for use in real time nucleic acid detection processes. Such real time nucleic acid detection processes are carried out with energy transfer elements attached to nucleic acid primers, nucleotides, nucleic acid probes or nucleic acid binding agents. Real time nucleic acid detection allows for the qualitative or quantitative detection or determination of single-stranded or double-stranded nucleic acids of interest in a sample. Other processes are provided by this invention including processes for removing a portion of a homopolymeric sequence, e.g., poly A sequence or tail, from an analyte or library of analytes. Compositions useful in carrying out such removal processes are also described and provided.

## CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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L16 ANSWER 22 OF 39 USPATFULL on STN
```

AN 2005:240503 USPATFULL

TΙ Chemical ligation of nucleic acids TN

Yowanto, Handy, Walnut, CA, UNITED STATES Yu, Changjun, Pasadena, CA, UNITED STATES ΡI US 20050208503 A1 20050922

A1 20040316 (10) US 2004-803166 ΑI

DT Utility

APPLICATION FS

LREP DORSEY & WHITNEY LLP, 555 CALIFORNIA STREET, SUITE 1000, SUITE 1000, SAN FRANCISCO, CA, 94104, US

Number of Claims: 30 CLMN Exemplary Claim: 1

14 Drawing Page(s)

LN.CNT 1735

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The invention relates to the field of nucleic acid analysis. More particularly, the invention relates to compositions and methods used for the detection of sequence variations or single nucleotide

```
L16 ANSWER 23 OF 39 USPATFULL on STN
AN
      2005:159178 USPATFULL
TT
      Real-time nucleic acid detection processes and compositions
      Rabbani, Elazar, New York, NY, UNITED STATES
      Stavrianopoulos, Jannis G., Bavsnore, NY, UNITED STATES
      Donegan, James J., Long Beach, NY, UNITED STATES
      Coleman, Jack, East Northport, NY, UNITED STATES
      Liu, Dakai, Islip, NY, UNITED STATES
PΙ
      US 20050137388
                       A1 20050623
ΑI
      US 2002-96076
                          A1 20020312 (10)
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DT Utility FS APPLICATION

LREP ENZO BIOCHEM, INC., 527 MADISON AVENUE (9TH FLOOR), NEW YORK, NY, 10022,

CLMN Number of Claims: 542 ECL Exemplary Claim: 1 DRWN 15 Drawing Page(s)

LN.CNT 6158

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB This invention provides for compositions for use in real time nucleic acid detection processes. Such real time nucleic acid detection processes are carried out with energy transfer elements attached to nucleic acid primers, nucleotides, nucleic acid probes or nucleic acid binding agents. Real time nucleic acid detection allows for the qualitative or quantitative detection or determination of single-stranded or double-stranded nucleic acids of interest in a sample. Other processes are provided by this invention including processes for removing a portion of a homopolymeric sequence, e.g., poly A sequence or tail, from an analyte or library of analytes. Compositions useful in carrying out such removal processes are also described and provided.

## CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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L16 ANSWER 24 OF 39 USPATFULL on STN
AN 2005:125227 USPATFULL
```

TI Mobility-modifying cyanine dyes

IN Menchen, Steven M., Fremont, CA, UNITED STATES Benson, Scott C., Alameda, CA, UNITED STATES Rosenblum, Barnett B., San Jose, CA, UNITED STATES Khan, Shaheer H., Foster City, CA, UNITED STATES

PA Applera Corporation, Foster City, CA, UNITED STATES (U.S. corporation)
PI US 20050107617 A1 20050519

PI US 20050107617 A1 20050519 AI US 2004-801092 A1 20040315 (10)

RLI Division of Ser. No. US 2000-477270, filed on 4 Jan 2000, GRANTED, Pat. No. US 6716994

DT Utility FS APPLICATION

LREP MILA KASAN, PATENT DEPT., APPLIED BIOSYSTEMS, 850 LINCOLN CENTRE DRIVE, FOSTER CITY, CA, 94404, US

CLMN Number of Claims: 41 ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 3875

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The present invention provides a novel class of fluorescent cyanine dye compounds that are modified at one of the hetercyclic ring nitrogen

atoms with a mobility-modifying moiety that permits the electrophoretic mobilities of polynucleotides labeled with the mobility-modifying cvanine dves to be adjusted or tuned in a predictable fashion while retaining enzymatic activity. The ability to predictably tune the relative electrophoretic mobilities of the dyes permits the creation of sets of mobility-matched fluorescent dyes of a variety of structures for a variety of applications, including fluorescence-based 4-color nucleic

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acid sequencing reactions.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L16 ANSWER 25 OF 39 USPATFULL on STN
AN
       2005:5243 USPATFULL
TT
       Novel chemiluminescent reagents
TN
       Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES
       Rabbani, Elazar, New York, NY, UNITED STATES
       Enzo Life Sciences, Inc., New York, NY, 10022 (U.S. corporation)
PΑ
PΙ
      US 20050004350
                          A1 20050106
      US 7256299
                          B2 20070814
      US 2004-764388
                          A1
                              20040123 (10)
AΙ
       Division of Ser. No. US 2002-96075, filed on 12 Mar 2002, PENDING
RLI
DT
      Utility
FS
      APPLICATION
LREP
      Ronald C. Fedus, Esq., Enzo Life Sciences, Inc., c/o Enzo Biochem, Inc.,
      527 Madison Avenue (9th Floor), New York, NY, 10022-4304
      Number of Claims: 17
CLMN
      Exemplary Claim: CLM-1-286
ECL
DRWN
      15 Drawing Page(s)
LN.CNT 3601
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
      This invention provides for labeling reagents, labeled targets and
       processes for preparing labeling reagents. The labeling reagents can
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take the form of cyanine dyes, xanthene dyes, porphyrin dyes, coumarin dyes or composite dyes. These labeling reagents are useful for labeling probes or targets, including nucleic acids and proteins. These reagents can be usefully applied to protein and nucleic acid probe based assays. They are also applicable to real-time detection processes.

# CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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L16 ANSWER 26 OF 39 USPATFULL on STN
AN
       2004:321700 USPATFULL
ΤI
       Labeling reagents comprising aphenylic analogs of rhodamine dyes
IN
       Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES
       Rabbani, Elazar, New York, NY, UNITED STATES
PA
       Enzo Life Sciences, Inc., New York, NY (U.S. corporation)
ΡI
      US 20040254355
                          A1 20041216
      US 7256291
                          B2 20070814
      US 2004-763076
ΑI
                         A1 20040122 (10)
RLI
      Division of Ser. No. US 2002-96075, filed on 12 Mar 2002, PENDING
DT
      Utility
FS
      APPLICATION
       Ronald C. Fedus, Esq., Enzo Life Sciences, Inc., c/o Enzo Biochem, Inc.,
LREP
       527 Madison Avenue (9th Floor), New York, NY, 10022-4304
      Number of Claims: 286
CLMN
ECL
      Exemplary Claim: 1
DRWN
      15 Drawing Page(s)
LN.CNT 4545
```

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB This invention provides for labeling reagents, labeled targets and processes for preparing labeling reagents. The labeling reagents can take the form of cyanine dyes, xanthene dyes, porphyrin dyes, coumarin dyes or composite dyes. These labeling reagents are useful for labeling probes or targets, including nucleic acids and proteins. These reagents can be usefully applied to protein and nucleic acid probe based assays. They are also applicable to real-time detection processes.

# CAS INDEXING IS AVAILABLE FOR THIS PATENT.

- L16 ANSWER 27 OF 39 USPATFULL on STN
- AN 2004:292946 USPATFULL
- ΤI Heterodimeric dye composition
- Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES IN
  - Rabban, Elazar, New York, NY, UNITED STATES
- PA Enzo Life Sciences, Inc., New York, NY, UNITED STATES, 10022 (U.S.
- corporation) PΤ US 20040230036 A1 20041118
- US 7323571 B2 20080129 AΙ US 2004-764389 A1 20040123 (10)
- RLI
- Division of Ser. No. US 2002-96075, filed on 12 Mar 2002, PENDING DT Utility
- FS APPLICATION
- LREP Ronald C. Fedus, Esq., Enzo Life Sciences, Inc., c/o Enzo Biochem, Inc., 527 Madison Avenue (9th Floor), New York, NY, 10022-4304
- CLMN Number of Claims: 286
- Exemplary Claim: 1
- 15 Drawing Page(s)
- LN.CNT 4541
- CAS INDEXING IS AVAILABLE FOR THIS PATENT.
- This invention provides for labeling reagents, labeled targets and processes for preparing labeling reagents. The labeling reagents can take the form of cyanine dyes, xanthene dyes, porphyrin dyes, coumarin dyes or composite dyes. These labeling reagents are useful for labeling probes or targets, including nucleic acids and proteins. These reagents can be usefully applied to protein and nucleic acid probe based assays. They are also applicable to real-time detection processes.

## CAS INDEXING IS AVAILABLE FOR THIS PATENT.

- L16 ANSWER 28 OF 39 USPATFULL on STN AN 2004:292164 USPATFULL
  - Novel dve labeling composition
  - Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES IN
  - Rabbani, Elazar, New York, NY, UNITED STATES
  - Enzo Life Sciences, Inc., New York, NY, 10022 (U.S. corporation) PA PΙ US 20040229248 A1 20041118
  - B2 20050927 US 6949659
  - US 2004-764393 A1 20040123 (10) ΑI
  - Division of Ser. No. US 2002-96075, filed on 12 Mar 2002, PENDING RLI DT Utility
  - FS APPLICATION
  - LREP Ronald C. Fedus, Esq., Enzo Life Sciences, Inc., c/o Enzo Biochem, Inc., 527 Madison Avenue, 9th Floor, New York, NY, 10022-4304
- CLMN Number of Claims: 4 ECL Exemplary Claim: CLM-1-286
- 15 Drawing Page(s)
- LN.CNT 3537
- CAS INDEXING IS AVAILABLE FOR THIS PATENT.
- This invention provides for labeling reagents, labeled targets and processes for preparing labeling reagents. The labeling reagents can take the form of cyanine dyes, xanthene dyes, porphyrin dyes, coumarin dyes or composite dyes. These labeling reagents are useful for labeling

probes or targets, including nucleic acids and proteins. These reagents can be usefully applied to protein and nucleic acid probe based assays. They are also applicable to real-time detection processes.

## CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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L16 ANSWER 29 OF 39 USPATFULL on STN
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AN 2004:260541 USPATFULL TI Process for preparing n

Process for preparing novel cyanine dye labeling reagents

IN Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES

Rabbam, Elazar, New York, NY, UNITED STATES

PA Enzo Life Sciences, Inc., New York, NY, 10022 (U.S. corporation)

PI US 20040203038 A1 20041014 US 7241897 B2 20070710

AI US 2004-761906 A1 20040121 (10)

RLI Division of Ser. No. US 2002-96075, filed on 12 Mar 2002, PENDING

DT Utility FS APPLICAT

FS APPLICATION
LREP Ronald C. Fedus, Esq., Enzo Life Sciences, Inc., c/o Enzo Biochem, Inc.,
527 Madison Avenue (9th Floor), New York, NY, 10022-4304

CLMN Number of Claims: 15

ECL Exemplary Claim: CLM-1-286 DRWN 15 Drawing Page(s)

LN.CNT 3584

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

This invention provides for labeling reagents, labeled targets and processes for preparing labeling reagents. The labeling reagents can take the form of cyanine dyes, xanthene dyes, porphyrin dyes, coumarin dyes or composite dyes. These labeling reagents are useful for labeling probes or targets, including nucleic acids and proteins. These reagents can be usefully applied to protein and nucleic acid probe based assays. They are also applicable to real-time detection processes.

# CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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L16 ANSWER 30 OF 39 USPATFULL on STN
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AN 2004:248291 USPATFULL

 $\ensuremath{\mathsf{TI}}$  Process for detecting the presence or quantity of enzymatic activity in a sample

N Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES Rabbani, Elazar, New York, NY, UNITED STATES

PA Enzo Life Sciences, Inc., New York, NY, UNITED STATES, 10022 (U.S.

corporation)
PI US 20040192893 A1 20040930
US 7553959 B2 20090630

AI US 2004-764417 A1 20040123 (10)
RLI Division of Ser. No. US 2002-96075, filed on 12 Mar 2002, PENDING

DIVISION DIVISION

LREP

DT Utility
FS APPLICATION

Ronald C. Fedus, Esq., Enzo Life Sciences, Inc., c/o Enzo Biochem, Inc.,

527 Madison Avenue (9th Floor), New York, NY, 10022-4304 CLMN Number of Claims: 36

ECL Exemplary Claim: CLM-1-286

DRWN 15 Drawing Page(s) LN.CNT 3665

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB This invention provides for labeling reagents, labeled targets and processes for preparing labeling reagents. The labeling reagents can take the form of cyanine dyes, xanthene dyes, porphyrin dyes, coumarin dyes or composite dyes. These labeling reagents are useful for labeling probes or targets, including nucleic acids and proteins. These reagents

can be usefully applied to protein and nucleic acid probe based assays. They are also applicable to real-time detection processes.

# CAS INDEXING IS AVAILABLE FOR THIS PATENT.

imidazole-PEG compounds.

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L16 ANSWER 31 OF 39 USPATFULL on STN
AN
       2004:228200 USPATFULL
       Process for detecting the presence or quantity of enzymatic activity in
       a sample
       Stavrianopoulos, Jannis G., Bayshore, NY, UNITED STATES
       Rabbani, Elazar, New York, NY, UNITED STATES
PA
       Enzo Life Sciences, Inc., New York, NY, UNITED STATES (U.S. corporation)
PΙ
       US 20040176586
                          A1 20040909
       US 7163796
                          B2 20070116
ΑТ
       US 2004-764418
                          A1 20040123 (10)
RLT
       Division of Ser. No. US 2002-96075, filed on 12 Mar 2002, PENDING
DТ
       Utility
FS
       APPLICATION
LREP
       Ronald C. Fedus, Esq., Enzo Life Sciences, Inc., c/o Enzo Biochem, Inc.,
       527 Madison Avenue (9th Floor), New York, NY, 10022-4304
       Number of Claims: 286
CLMN
ECL
       Exemplary Claim: 1
DRWN
       15 Drawing Page(s)
LN.CNT 4543
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       This invention provides for labeling reagents, labeled targets and
       processes for preparing labeling reagents. The labeling reagents can
       take the form of cyanine dyes, xanthene dyes, porphyrin dyes, coumarin
       dyes or composite dyes. These labeling reagents are useful for labeling
       probes or targets, including nucleic acids and proteins. These reagents
       can be usefully applied to protein and nucleic acid probe based assays.
       They are also applicable to real-time detection processes.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L16 ANSWER 32 OF 39 USPATFULL on STN
AN
       2004:221271 USPATFULL
ΤI
       Detectable labeled nucleoside analogs and methods of use
       Bodepudi, Veeraiah, San Ramon, CA, UNITED STATES
       Gupta, Amar, Danville, CA, UNITED STATES
       Will, Stephen, Oakland, CA, UNITED STATES
PT
       US 20040171040
                          A1 20040902
       US 7220847
                          B2 20070522
AΙ
       US 2003-719257
                          A1 20031121 (10)
PRAT
       US 2002-428484P
                               20021122 (60)
DT
       Utility
FS
       APPLICATION
LREP
      MORGAN, LEWIS & BOCKIUS, LLP., 3300 HILLVIEW AVENUE, PALO ALTO, CA,
       94304
CLMN
       Number of Claims: 80
ECL
       Exemplary Claim: 1
DRWN
      No Drawings
LN.CNT 2435
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       The invention relates to detectable labels useful for detection of
       nucleotide sequences. Specifically, the invention relates to
       labeled-imidazole-PEG compounds, such as nucleosides, nucleotides, and
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nucleic acids incorporating such compounds, and methods utilizing such compounds. The invention further relates to kits comprising labeled

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L16 ANSWER 33 OF 39 USPATFULL on STN
       2004:126981 USPATFULL
AN
TI
       Manipulation of microparticles in microfluidic systems
       Burd Mehta, Tammy, San Jose, CA, UNITED STATES
       Kopf-Sill, Anne R., Portola Valley, CA, UNITED STATES
       Parce, J. Wallace, Palo Alto, CA, UNITED STATES
       Chow, Andrea W., Los Altos, CA, UNITED STATES
       Bousse, Luc J., Los Altos, CA, UNITED STATES
       Knapp, Michael R., Redwood City, CA, UNITED STATES
       Nikiforov, Theo T., San Jose, CA, UNITED STATES
       Gallagher, Steve, Palo Alto, CA, UNITED STATES
PA
       Caliper Technologies Corp., Mountain View, CA (U.S. corporation)
PΤ
       US 20040096960
                          A1 20040520
ΑI
      US 2003-606201
                           A1 20030625 (10)
RLI
       Continuation of Ser. No. US 2000-510626, filed on 22 Feb 2000, GRANTED,
       Pat. No. US 6632655
PRAI
      US 1999-121223P
                               19990223 (60)
      US 1999-127825P
                               19990405 (60)
      US 1999-128643P
                               19990409 (60)
DT
      Utility
FS
       APPLICATION
LREP
       CALIPER LIFE SCIENCES, INC., 605 FAIRCHILD DRIVE, MOUNTAIN VIEW, CA,
       94043-2234
CLMN
      Number of Claims: 1
       Exemplary Claim: 1
ECL
DRWN
       19 Drawing Page(s)
LN.CNT 4120
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AB
       Arrays of flowable or fixed particle sets are used in microfluidic
       systems for performing assays and modifying hydrodynamic flow. Also
       provided are assays utilizing flowable or fixed particle sets within a
      microfluidic system, as well as kits, apparatus and integrated systems
      comprising arrays and array members.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L16 ANSWER 34 OF 39 USPATFULL on STN
AN
       2004:85289 USPATFULL
ΤI
       Mobility-Modifying Cyanine Dyes
IN
      Menchen, Steven M., Fremont, CA, United States
       Benson, Scott C., Alameda, CA, United States
       Rosenblum, Barnett B., San Jose, CA, United States
       Khan, Shaheer H., Foster City, CA, United States
       Applera Corporation, Foster City, CA, United States (U.S. corporation)
PA
ΡI
      US 6716994
                          B1 20040406
      US 2000-477270
ΑI
                               20000104 (9)
DT
      Utility
FS
      GRANTED
EXNAM Primary Examiner: McKane, Joseph K.; Assistant Examiner: Saeed, Kamal
LREP
       Peasu, Ann, Liptak, Vincent P.
CLMN
      Number of Claims: 5
       Exemplary Claim: 1
      0 Drawing Figure(s); 0 Drawing Page(s)
LN.CNT 3306
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AB
       The present invention provides a novel class of fluorescent cyanine dye
       compounds that are modified at one of the hetercyclic ring nitrogen
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atoms with a mobility-modifying moiety that permits the electrophoretic

mobilities of polynucleotides labeled with the mobility-modifying cyanine dyes to be adjusted or tuned in a predictable fashion while retaining enzymatic activity. The ability to predictably tune the relative electrophoretic mobilities of the dyes permits the creation of sets of mobility-matched fluorescent dyes of a variety of structures for a variety of applications, including fluorescence-based 4-color nucleic acid sequencing reactions.

# CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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L16 ANSWER 35 OF 39 USPATFULL on STN
AN
       2004:44533 USPATFULL
TI
       Releasable nonvolatile mass-label molecules
TN
       Monforte, Joseph A., Berkeley, CA, UNITED STATES
       Becker, Christopher H., Palo Alto, CA, UNITED STATES
       Pollart, Daniel J., Menlo Park, CA, UNITED STATES
       Shaler, Thomas A., Menlo Park, CA, UNITED STATES
                          A1 20040219
PΙ
       US 20040033525
       US 2003-637935
ΑI
                           A1 20030807 (10)
RLI
       Division of Ser. No. US 2002-202189, filed on 22 Jul 2002, PENDING
       Continuation of Ser. No. US 1997-988024, filed on 10 Dec 1997, GRANTED,
       Pat. No. US 6635452
      US 1996-33037P
PRAI
                               19961210 (60)
      US 1997-46719P
                               19970516 (60)
       Utility
       APPLICATION
LREP
       HELLER EHRMAN WHITE & MCAULIFFE LLP, 4350 LA JOLLA VILLAGE DRIVE, 7TH
       FLOOR, SAN DIEGO, CA, 92122-1246
CLMN
      Number of Claims: 58
ECL
      Exemplary Claim: 1
DRWN
      35 Drawing Page(s)
LN.CNT 3933
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
```

AB Releasable tag reagents for use in the detection and analysis of target molecules, particular in mass spectrometric analyses are provided. Also provided are methods of detection that employ releasable tag reagents.

Exemplary Claim: 1

4 Drawing Figure(s); 4 Drawing Page(s)

ECI.

DRWN

LN.CNT 1299

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CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L16 ANSWER 36 OF 39 USPATFULL on STN
AN
       1999:110164 USPATFULL
       Ligase/polymerase-mediated genetic bit analysis of single
TI
       nucleotide polymorphisms and its use in genetic analysis
IN
       Nikiforov, Theo, Baltimore, MD, United States
       Karn, Jonathan, Little Shelord, United Kingdom
       Goelet, Philip, Cockeysville, MD, United States
      Orchid Biocomputer, Inc., Princeton, NJ, United States (U.S.
PA
       corporation)
      US 5952174
PΤ
                               19990914
      US 1997-929101
AΙ
                               19970915 (8)
RLI
       Continuation of Ser. No. US 1996-694835, filed on 9 Aug 1996, now
       patented, Pat. No. US 5679524 which is a continuation of Ser. No. US
       1994-192631, filed on 7 Feb 1994, now abandoned
DT
      Utility
FS
       Granted
EXNAM Primary Examiner: Marschel, Ardin H.
LREP
      Auerbach, Jeffrey I., Mendelson, Elliot C.Howrey & Simon
CLMN
      Number of Claims: 13
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#### CAS INDEXING IS AVAILABLE FOR THIS PATENT. A method is provided for determining the identity of a AB nucleotide at a preselected site in a nucleic acid molecule. The method involves the incorporation of a nucleoside triphosphate that is complementary to the nucleotide present at the preselected site onto the terminus of a primer molecule, and their subsequent ligation to a second oligonucleotide. The reaction is monitored by detecting a specific label attached to the reaction's solid phase or by detection in solution. CAS INDEXING IS AVAILABLE FOR THIS PATENT. L16 ANSWER 37 OF 39 USPATFULL on STN AN 97:96725 USPATFULL ΤТ Ligase/polymerase mediated genetic bit analysis of single nucleotide polymorphisms and its use in genetic analysis Nikiforov, Theo, Baltimore, MD, United States TN Karn, Jonathan, Little Shelord, United Kingdom Goelet, Philip, Cockeysville, MD, United States PA Molecular Tool, Inc., Baltimore, MD, United States (U.S. corporation) ΡI US 5679524 19971021 ΑI US 1996-694835 19960809 (8) Continuation of Ser. No. US 1994-192631, filed on 7 Feb 1994, now RLI abandoned Utility FS Granted EXNAM Primary Examiner: Marschel, Ardin H. LREP Howrev & Simon, Auerbach, Jeffrev I. CLMN Number of Claims: 24 ECL Exemplary Claim: 1 DRWN 4 Drawing Figure(s); 4 Drawing Page(s) LN.CNT 1456 CAS INDEXING IS AVAILABLE FOR THIS PATENT. AB A method is provided for determining the identity of a nucleotide at a preselected site in a nucleic acid molecule. The method involves the incorporation of a nucleoside triphosphate that is complementary to the nucleotide present at the preselected site onto the terminus of a primer molecule, and their subsequent ligation to a second oligonucleotide. The reaction is monitored by detecting a specific label attached to the reaction's solid phase or by detection in solution. CAS INDEXING IS AVAILABLE FOR THIS PATENT. L16 ANSWER 38 OF 39 USPATFULL on STN AN 92:80906 USPATFULL ΤТ Alkvnvlamino-nucleotides IN Hobbs, Jr., Frank W., Wilmington, DE, United States Trainor, George L., Wilmington, DE, United States PA E. I. Du Pont de Nemours and Company, Wilmington, DE, United States (U.S. corporation) PΙ US 5151507 19920929 US 1991-713906 ΑI 19910612 (7) 20080910 DCD

Continuation-in-part of Ser. No. US 1987-57565, filed on 12 Jun 1987,

now patented, Pat. No. US 5047519 which is a continuation-in-part of Ser. No. US 1986-881372, filed on 2 Jul 1986, now abandoned

EXNAM Primary Examiner: Brown, Johnnie R.; Assistant Examiner: Wilson, J.

RLI

Utility Granted

Oliver

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LREP Frank, George A.
CLMN Number of Claims: 9
ECL.
      Exemplary Claim: 1
DRWN No Drawings
LN.CNT 3011
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       Alkynylamino-nucleotides and labeled alkynylaminonucleotides useful, for
       example, as chain terminating substrates for DNA sequencing are provided
       along with several key intermediates and processes for their
       preparation. For some applications, longer, hydrophilic linkers are
       provided.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L16 ANSWER 39 OF 39 USPATFULL on STN
       91:73471 USPATFULL
AN
тт
       Alkynylamino-nucleotides
IN
       Hobbs, Jr., Frank W., Wilmington, DE, United States
       Cocuzza, Anthony J., Wilmington, DE, United States
PA
       E. I. Du Pont de Nemours and Company, Wilmington, DE, United States
       (U.S. corporation)
PΤ
       US 5047519
                               19910910
       US 1987-57565
ΑI
                               19870612 (7)
RLI
       Continuation-in-part of Ser. No. US 1986-881372, filed on 2 Jul 1986,
       now abandoned
       Utility
FS
       Granted
EXNAM Primary Examiner: Brown, Johnnie R.; Assistant Examiner: Wilson, James
LREP
      Frank, George A.
CLMN Number of Claims: 12
ECL
       Exemplary Claim: 6,9
DRWN
      No Drawings
```

LN.CNT 2907

AB Alkynylamino-nucleotides and labeled alkynylamino-nucleotides useful, for example, as chain terminating substrates for DNA sequencing are provided along with several key intermediates and processes for their preparation.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.